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(54) Title: HIGH MOLECULAR WEIGHT SURFACE PROTEINES OF NON-TYPEABLE HAEMOPHILUS (57) Abstract High molecular weight surface proteins of non-typeable <i>Haemophilus influenzae</i> which exhibit immunogenic properties and genes encoding the same are described. Specifically, genes coding for two immunodominant high molecular weight proteins, HMW1 and HMW2, have been cloned, expressed and sequenced, while genes coding for high molecular proteins HMW3 and HMW4 have been cloned, expressed and partially sequenced.		

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TITLE OF INVENTIONHIGH MOLECULAR WEIGHT SURFACE PROTEINS
OF NON-TYPEABLE HAEMOPHILUSFIELD OF INVENTION

5 This invention relates to high molecular weight proteins of non-typeable haemophilus.

BACKGROUND TO THE INVENTION

10 Non-typeable Haemophilus influenzae are non-encapsulated organisms that are defined by their lack of reactivity with antisera against known H. influenzae capsular antigens.

15 These organisms commonly inhabit the upper respiratory tract of humans and are frequently responsible for infections, such as otitis media, sinusitis, conjunctivitis, bronchitis and pneumonia. Since these organisms do not have a polysaccharide capsule, they are not controlled by the present Haemophilus influenzae type b (Hib) vaccines, which are directed towards Hib bacterial capsular polysaccharides. 20 The non-typeable strains, however, do produce surface antigens that can elicit bactericidal antibodies. Two of the major outer membrane proteins, P2 and P6, have been identified as targets of human serum bactericidal activity. However, it has been shown that the P2 protein 25 sequence is variable, in particular in the non-typeable Haemophilus strains. Thus, a P2-based vaccine would not protect against all strains of the organism.

30 There have previously been identified by Barenkamp et al (Pediatr. Infect. Dis. J., 9:333-339, 1990) a group of high-molecular-weight (HMW) proteins that appeared to be major targets of antibodies present in human convalescent sera. Examination of a series of middle ear isolates revealed the presence of one or two such proteins in most strains. However, prior to the present 35 invention, the structures of these proteins were unknown as were pure isolates of such proteins.

SUMMARY OF INVENTION

The inventors, in an effort to further characterize the high molecular weight (HMW) Haemophilus proteins, have cloned, expressed and sequenced the genes coding for two immunodominant HMW proteins (designated HMW1 and HMW2) from a prototype non-typeable Haemophilus strain and have cloned, expressed and almost completely sequenced the genes coding for two additional immunodominant HMW proteins (designated HMW3 and HMW4) from another non-typeable Haemophilus strain.

In accordance with one aspect of the present invention, therefore, there is provided an isolated and purified gene coding for a high molecular weight protein of a non-typeable Haemophilus strain, particularly a gene coding for protein HMW1, HMW2, HMW3 or HMW4, as well as any variant or fragment of such protein which retains the immunological ability to protect against disease caused by a non-typeable Haemophilus strain. In another aspect, the invention provides a high molecular weight protein of non-typeable Haemophilus influenzae which is encoded by these genes.

BRIEF DESCRIPTION OF DRAWINGS

Figure 1 is a DNA sequence of a gene coding for protein HMW1 (SEQ ID NO: 1);

Figure 2 is a derived amino acid sequence of protein HMW1 (SEQ ID NO: 2);

Figure 3 is a DNA sequence of a gene coding for protein HMW2 (SEQ ID NO: 3);

Figure 4 is a derived amino acid sequence of HMW2 (SEQ ID NO: 4);

Figure 5A shows restriction maps of representative recombinant phages which contained the HMW1 or HMW2 structural genes, the locations of the structural genes being indicated by the shaded bars;

Figure 5B shows the restriction map of the T7 expression vector pT7-7;

Figure 6 contains the DNA sequence of a gene cluster for the hmw1 gene (SEQ ID NO: 5), comprising nucleotides 351 to 4958 (ORF a) (as in Figure 1), as well as two additional downstream genes in the 3' flanking region, comprising ORFs b, nucleotides 5114-6748 and c nucleotides 7062-9011;

Figure 7 contains the DNA sequence of a gene cluster for the hmw2 gene (SEQ ID NO: 6), comprising nucleotides 792 to 5222 (ORF a) (as in Figure 3), as well as two additional downstream genes in the 3' flanking region, comprising ORFs b, nucleotides 5375-7009, and c, nucleotides 7249-9198;

Figure 8 is a partial DNA sequence of a gene coding for protein HMW3 (SEQ ID NO: 7);

Figure 9 is a partial DNA sequence of a gene coding for protein HMW4 (SEQ ID NO: 8); and

Figure 10 is a comparison table for the derived amino acid sequence for proteins HMW1, HMW2, HMW3 and HMW4.

GENERAL DESCRIPTION OF INVENTION

The DNA sequences of the genes coding for HMW1 and HMW2, shown in Figures 1 and 3 respectively, were shown to be about 80% identical, with the first 1259 base pairs of the genes being identical. The derived amino acid sequences of the two HMW proteins, shown in Figures 2 and 4 respectively, are about 70% identical. Furthermore, the encoded proteins are antigenically related to the filamentous hemagglutinin surface protein of Bordetella pertussis. A monoclonal antibody prepared against filamentous hemagglutinin (FHA) of Bordetella pertussis was found to recognize both of the high molecular weight proteins. This data suggests that the HMW and FHA proteins may serve similar biological functions. The derived amino acid sequences of the HMW1 and HMW2 proteins show sequence similarity to that for the FHA protein. It has further been shown that these

antigenically-related proteins are produced by the majority of the non-typeable strains of Haemophilus. Antisera raised against the protein expressed by the HMW1 gene recognizes both the HMW2 protein and the B. pertussis FHA. The present invention includes an isolated and purified high molecular weight protein of non-typeable haemophilus which is antigenically related to the B. pertussis FHA, which may be obtained from natural sources or produced recombinantly.

10 A phage genomic library of a known strain of non-typeable Haemophilus was prepared by standard methods and the library was screened for clones expressing high molecular weight proteins, using a high titre antiserum against HMW's. A number of strongly reactive DNA clones
15 were plaque-purified and sub-cloned into a T7 expression plasmid. It was found that they all expressed either one or the other of the two high-molecular-weight proteins designated HMW1 and HMW2, with apparent molecular weights of 125 and 120 kDa, respectively, encoded by open reading
20 frames of 4.6 kb and 4.4 kb, respectively.

Representative clones expressing either HMW1 or HMW2 were further characterized and the genes isolated, purified and sequenced. The DNA sequence of HMW1 is shown in Figure 1 and the corresponding derived amino
25 acid sequence in Figure 2. Similarly, the DNA sequence of HMW2 is shown in Figure 3 and the corresponding derived amino acid sequence in Figure 4. Partial purification of the isolated proteins and N-terminal sequence analysis indicated that the expressed proteins are truncated since
30 their sequence starts at residue number 442 of both full length HMW1 and HMW2 gene products.

Subcloning studies with respect to the hmw1 and hmw2 genes indicated that correct processing of the HMW proteins required the products of additional downstream
35 genes. It has been found that both the hmw1 and hmw2 genes are flanked by two additional downstream open

reading frames (ORFs), designated b and c, respectively, (see Figures 6 and 7).

5 The b ORFs are 1635 bp in length, extending from nucleotides 5114 to 6748 in the case of hmw1 and nucleotides 5375 to 7009 in the case of hmw2, with their derived amino acid sequences 99% identical. The derived amino acid sequences demonstrate similarity with the derived amino acid sequences of two genes which encode proteins required for secretion and activation of hemolysins of P. mirabilis and S. marcescens.
10

The c ORFs are 1950 bp in length, extending from nucleotides 7062 to 9011 in the case of hmw1 and nucleotides 7249 to 9198 in the case of hmw2, with their derived amino acid sequences 96% identical. The hmw1 c ORF is preceded by a series of 9 bp direct tandem repeats. In plasmid subclones, interruption of the hmw1 b or c ORF results in defective processing and secretion of the hmw1 structural gene product.
15

The two high molecular weight proteins have been isolated and purified and shown to be partially protective against otitis media in chinchillas and to function as adhesins. These results indicate the potential for use of such high molecular proteins and structurally-related proteins of other non-typeable strains of Haemophilus influenzae as components in non-typeable Haemophilus influenzae vaccines.
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Since the proteins provided herein are good cross-reactive antigens and are present in the majority of non-typeable Haemophilus strains, it is evident that these HMW proteins may become integral constituents of a universal Haemophilus vaccine. Indeed, these proteins may be used not only as protective antigens against otitis, sinusitis and bronchitis caused by the non-typeable Haemophilus strains, but also may be used as carriers for the protective Hib polysaccharides in a conjugate vaccine against meningitis. The proteins also
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may be used as carriers for other antigens, haptens and polysaccharides from other organisms, so as to induce immunity to such antigens, haptens and polysaccharides.

5 The nucleotide sequences encoding two high molecular weight proteins of a different non-typeable Haemophilus strain (designated HMW3 and HMW4) have been largely elucidated, and are presented in Figures 8 and 9. HMW3 has an apparent molecular weight of 125 kDa while HMW4 has an apparent molecular weight of 123 kDa. These high
10 molecular weight proteins are antigenically related to the HMW1 and HMW2 proteins and to FHA. Sequence analysis of HMW3 is approximately 85% complete and of HMW4 95% complete, with short stretches at the 5'-ends of each gene remaining to be sequenced.

15 Figure 10 contains a multiple sequence comparison of the derived amino acid sequences for the four high molecular weight proteins identified herein. As may be seen from this comparison, stretches of identical peptide sequence may be found throughout the length of the
20 comparison, with HMW3 more closely resembling HMW1 and HMW4 more closely resembling HMW2. This information is highly suggestive of a considerable sequence homology between high molecular weight proteins from various non-typeable Haemophilus strains.

25 In addition, mutants of non-typeable H. influenzae strains that are deficient in expression of HMW1 or HMW2 or both have been constructed and examined for their capacity to adhere to cultured human epithelial cells. The hmw1 and hmw2 gene clusters have been expressed in E. coli and have been examined for in vitro adherence. The
30 results of such experimentation demonstrate that both HMW1 and HMW2 mediate attachment and hence are adhesins and that this function is present even in the absence of other H. influenzae surface structures.

35 With the isolation and purification of the high molecular weight proteins, the inventors are able to

determine the major protective epitopes by conventional epitope mapping and synthesize peptides corresponding to these determinants to be incorporated in fully synthetic or recombinant vaccines. Accordingly, the invention also comprises a synthetic peptide having an amino acid sequence corresponding to at least one protective epitope of a high molecular weight protein of a non-typeable Haemophilus influenzae. Such peptides are of varying length that constitute portions of the high-molecular-weight proteins, that can be used to induce immunity, either directly or as part of a conjugate, against the relative organisms and thus constitute vaccines for protection against the corresponding diseases.

The present invention also provides any variant or fragment of the proteins that retains the potential immunological ability to protect against disease caused by non-typeable Haemophilus strains. The variants may be constructed by partial deletions or mutations of the genes and expression of the resulting modified genes to give the protein variations.

EXAMPLES

Example 1:

Non-typeable H.influenzae strains 5 and 12 were isolated in pure culture from the middle ear fluid of children with acute otitis media. Chromosomal DNA from strain 12, providing genes encoding proteins HMW1 and HMW2, was prepared by preparing Sau3A partial restriction digests of chromosomal DNA and fractionating on sucrose gradients. Fractions containing DNA fragments in the 9 to 20 kbp range were pooled and a library was prepared by ligation into λ EMBL3 arms. Ligation mixtures were packaged in vitro and plate-amplified in a P2 lysogen of E. coli LE392.

For plasmid subcloning studies, DNA from a representative recombinant phage was subcloned into the

T7 expression plasmid pT7-7, containing the T7 RNA polymerase promoter $\Phi 10$, a ribosome-binding site and the translational start site for the T7 gene 10 protein upstream from a multiple cloning site (see Figure 5B).

5 DNA sequence analysis was performed by the dideoxy method and both strands of the HMW1 gene and a single strand of the HMW2 gene were sequenced.

Western immunoblot analysis was performed to identify the recombinant proteins being produced by
10 reactive phage clones. Phage lysates grown in LE392 cells or plaques picked directly from a lawn of LE392 cells on YT plates were solubilized in gel electrophoresis sample buffer prior to electrophoresis. Sodium dodecyl sulfate (SDS)-polyacrylamide gel
15 electrophoresis was performed on 7.5% or 11% polyacrylamide modified Laemmli gels. After transfer of the proteins to nitrocellulose sheets, the sheets were probed sequentially with an E. coli-absorbed human serum sample containing high-titer antibody to the high-
20 molecular-weight proteins and then with alkaline phosphatase-conjugated goat anti-human immunoglobulin G (IgG) second antibody. Sera from healthy adults contains high-titer antibody directed against surface-exposed high-molecular-weight proteins of non-typeable H. influenzae.
25 One such serum sample was used as the screening antiserum after having been extensively absorbed with LE392 cells.

To identify recombinant proteins being produced by E. coli transformed with recombinant plasmids, the
30 plasmids of interest were used to transform E. coli BL21 (DE3)/pLysS. The transformed strains were grown to an A_{600} of 0.5 in L broth containing 50 μ g of ampicillin per ml. IPTG was then added to 1 mM. One hour later, cells were harvested, and a sonicate of the cells was prepared.
35 The protein concentrations of the samples were determined by the bicinchoninic acid method. Cell sonicates

containing 100 μ g of total protein were solubilized in electrophoresis sample buffer, subjected to SDS-polyacrylamide gel electrophoresis, and transferred to nitrocellulose. The nitrocellulose was then probed sequentially with the E. coli-absorbed adult serum sample and then with alkaline phosphatase-conjugated goat anti-human IgG second antibody.

Western immunoblot analysis also was performed to determine whether homologous and heterologous non-typeable H. influenzae strains expressed high-molecular-weight proteins antigenically related to the protein encoded by the cloned HMW1 gene (rHMW1). Cell sonicates of bacterial cells were solubilized in electrophoresis sample buffer, subjected to SDS-polyacrylamide gel electrophoresis, and transferred to nitrocellulose. Nitrocellulose was probed sequentially with polyclonal rabbit rHMW1 antiserum and then with alkaline phosphatase-conjugated goat anti-rabbit IgG second antibody.

Finally, Western immunoblot analysis was performed to determine whether non-typeable Haemophilus strains expressed proteins antigenically related to the filamentous hemagglutinin protein of Bordetella pertussis. Monoclonal antibody X3C, a murine immunoglobulin G (IgG) antibody which recognizes filamentous hemagglutinin, was used to probe cell sonicates by Western blot. An alkaline phosphatase-conjugated goat anti-mouse IgG second antibody was used for detection.

To generate recombinant protein antiserum, E. coli BL21(DE3)/pLySS was transformed with pHMW1-4, and expression of recombinant protein was induced with IPTG, as described above. A cell sonicate of the bacterial cells was prepared and separated into a supernatant and pellet fraction by centrifugation at 10,000 x g for 30 min. The recombinant protein fractionated with the

pellet fraction. A rabbit was subcutaneously immunized on biweekly schedule with 1 mg of protein from the pellet fraction, the first dose given with Freund's complete adjuvant and subsequent doses with Freund's incomplete adjuvant. Following the fourth injection, the rabbit was bled. Prior to use in the Western blot assay, the antiserum was absorbed extensively with sonicates of the host E. coli strain transformed with cloning vector alone.

To assess the sharing of antigenic determinants between HMW1 and filamentous hemagglutinin, enzyme-linked immunosorbent assay (ELISA) plates (Costar, Cambridge, Mass.) were coated with 60 μ l of a 4-ug/ml solution of filamentous hemagglutinin in Dulbecco's phosphate-buffered saline per well for 2 h at room temperature. Wells were blocked for 1 h with 1% bovine serum albumin in Dulbecco's phosphate-buffered saline prior to addition of serum dilutions. rHMW1 antiserum was serially diluted in 0.1% Brij (Sigma, St. Louis, Mo.) in Dulbecco's phosphate-buffered saline and incubated for 3 h at room temperature. After being washed, the plates were incubated with peroxidase-conjugated goat anti-rabbit IgG antibody (Bio-Rad) for 2 h at room temperature and subsequently developed with 2,2'-azino-bis(3-ethylbenzthiazoline-6-sulfonic acid) (Sigma) at a concentration of 0.54 in mg/ml in 0.1 M sodium citrate buffer, pH 4.2, containing 0.03% H₂O₂. Absorbances were read on an automated ELISA reader.

Recombinant phage expressing HMW1 or HMW2 were recovered as follows. The non-typeable H. influenzae strain 12 genomic library was screened for clones expressing high-molecular-weight proteins with an E. coli-absorbed human serum sample containing a high titer of antibodies directed against the high-molecular-weight proteins.

Numerous strongly reactive clones were identified along with more weakly reactive ones. Twenty strongly reactive clones were plaque-purified and examined by Western blot for expression of recombinant proteins. Each of the strongly reactive clones expressed one of two types of high-molecular-weight proteins, designated HMW1 and HMW2. The major immunoreactive protein bands in the HMW1 and HMW2 lysates migrated with apparent molecular masses of 125 and 120 kDa, respectively. In addition to the major bands, each lysate contained minor protein bands of higher apparent molecular weight. Protein bands seen in the HMW2 lysates at molecular masses of less than 120 kDa were not regularly observed and presumably represent proteolytic degradation products. Lysates of LE392 infected with the λ EMBL3 cloning vector alone were non-reactive when immunologically screened with the same serum sample. Thus, the observed activity was not due to cross-reactive E. coli proteins or λ EMBL3-encoded proteins. Furthermore, the recombinant proteins were not simply binding immunoglobulin nonspecifically, since the proteins were not reactive with the goat anti-human IgG conjugate alone, with normal rabbit sera, or with serum from a number of healthy young infants.

Representative clones expressing either the HMW1 or HMW2 recombinant proteins were characterized further. The restriction maps of the two phage types were different from each other, including the regions encoding the HMW1 and HMW2 structural genes. Figure 5A shows restriction maps of representative recombinant phage which contained the HMW1 or HMW2 structural genes. The locations of the structural genes are indicated by the shaded bars.

HMW1 plasmid subclones were constructed by using the T7 expression plasmid T7-7 (Fig. 5A and B). HMW2 plasmid subclones also were constructed, and the results with

these latter subclones were similar to those observed with the HMW1 constructs.

The approximate location and direction of transcription of the HMW1 structure gene were initially determined by using plasmid pHMW1 (Fig. 5A). This plasmid was constructed by inserting the 8.5-kb BamHI-SalI fragment from λ HMW1 into BamHI- and SalI-cut pT7-7. E. coli transformed with pHMW1 expressed an immunoreactive recombinant protein with an apparent molecular mass of 115 kDa, which was strongly inducible with IPTG. This protein was significantly smaller than the 125-kDa major protein expressed by the parent phage, indicating that it either was being expressed as a fusion protein or was truncated at the carboxy terminus.

To more precisely localize the 3' end of the structural gene, additional plasmids were constructed with progressive deletions from the 3' end of the pHMW1 construct. Plasmid pHMW1-1 was constructed by digestion of pHMW1 with PstI, isolation of the resulting 8.8-kb fragment, and religation. Plasmid pHMW1-2 was constructed by digestion of pHMW1 with HindIII, isolation of the resulting 7.5-kb fragment, and religation. E. coli transformed with either plasmid pHMW1-1 or pHMW1-2 also expressed an immunoreactive recombinant protein with an apparent molecular mass of 115 kDa. These results indicated that the 3' end of the structural gene was 5' of the HindIII site.

To more precisely localize the 5' end of the gene, plasmids pHMW1-4 and pHMW1-7 were constructed. Plasmid pHMW1-4 was constructed by cloning the 5.1-kb BamHI-HindIII fragment from λ HMW1 into a pT7-7-derived plasmid containing the upstream 3.8-kb EcoRI-BamHI fragment. E. coli transformed with pHMW1-4 expressed an immunoreactive protein with an apparent molecular mass of approximately 160 kDa. Although protein production was inducible with IPTG, the levels of protein production in these

transformants were substantially lower than those with the pHMW1-2 transformants described above. Plasmid pHMW1-7 was constructed by digesting pHMW1-4 with NdeI and SpeI. The 9.0-kbp fragment generated by this double digestion was isolated, blunt ended, and religated. E. coli transformed with pHMW1-7 also expressed an immunoreactive protein with an apparent molecular mass of 160 kDa, a protein identical in size to that expressed by the pHMW1-4 transformants. The result indicated that the initiation codon for the HMW1 structural gene was 3' of the SpeI site. DNA sequence analysis confirmed this conclusion.

As noted above, the λ HMW1 phage clones expressed a major immunoreactive band of 125 kDa, whereas the HMW1 plasmid clones pHMW1-4 and pHMW1-7, which contained what was believed to be the full-length gene, expressed an immunoreactive protein of approximately 160 kDa. This size discrepancy was disconcerting. One possible explanation was that an additional gene or genes necessary for correct processing of the HMW1 gene product were deleted in the process of subcloning. To address this possibility, plasmid pHMW1-14 was constructed. This construct was generated by digesting pHMW1 with NdeI and MluI and inserting the 7.6-kbp NdeI-MluI fragment isolated from pHMW1-4. Such a construct would contain the full-length HMW1 gene as well as the DNA 3' of the HMW1 gene which was present in the original HMW1 phage. E. coli transformed with this plasmid expressed major immunoreactive proteins with apparent molecular masses of 125 and 160 kDa as well as additional degradation products. The 125- and 160-kDa bands were identical to the major and minor immunoreactive bands detected in the HMW1 phage lysates. Interestingly, the pHMW1-14 construct also expressed significant amounts of protein in the uninduced condition, a situation not observed with the earlier constructs.

The relationship between the 125- and 160-kDa proteins remains somewhat unclear. Sequence analysis, described below, reveals that the HMW1 gene would be predicted to encode a protein of 159 kDa. It is believed
5 that the 160-kDa protein is a precursor form of the mature 125-kDa protein, with the conversion from one protein to the other being dependent on the products of the two downstream genes.

Sequence analysis of the HMW1 gene (Figure 1)
10 revealed a 4,608-bp open reading frame (ORF), beginning with an ATG codon at nucleotide 351 and ending with a TAG stop codon at nucleotide 4959. A putative ribosome-binding site with the sequence AGGAG begins 10 bp upstream of the putative initiation codon. Five other in-
15 frame ATG codons are located within 250 bp of the beginning of the ORF, but none of these is preceded by a typical ribosome-binding site. The 5'-flanking region of the ORF contains a series of direct tandem repeats, with the 7-bp sequence ATCTTTC repeated 16 times. These
20 tandem repeats stop 100 bp 5' of the putative initiation codon. An 8-bp inverted repeat characteristic of a rho-independent transcriptional terminator is present, beginning at nucleotide 4983, 25 bp 3' of the presumed translational stop. Multiple termination codons are
25 present in all three reading frames both upstream and downstream of the ORF. The derived amino acid sequence of the protein encoded by the HMW1 gene (Figure 2) has a molecular weight of 159,000, in good agreement with the apparent molecular weights of the proteins expressed by
30 the HMW1-4 and HMW1-7 transformants. The derived amino acid sequence of the amino terminus does not demonstrate the characteristics of a typical signal sequence. The BamHI site used in generation of pHMW1 comprises bp 1743 through 1748 of the nucleotide sequence. The ORF
35 downstream of the BamHI site would be predicted to encode a protein of 111 kDa, in good agreement with the 115 kDa

estimated for the apparent molecular mass of the pHMW1-encoded fusion protein.

The sequence of the HMW2 gene (Figure 3) consists of a 4,431-bp ORF, beginning with an ATG codon at nucleotide 352 and ending with a TAG stop codon at nucleotide 4783. The first 1,259 bp of the ORF of the HMW2 gene are identical to those of the HMW1 gene. Thereafter, the sequences begin to diverge but are 80% identical overall. With the exception of a single base addition at nucleotide 93 of the HMW2 sequence, the 5'-flanking regions of the HMW1 and HMW2 genes are identical for 310 bp upstream from the respective initiation codons. Thus, the HMW2 gene is preceded by the same set of tandem repeats and the same putative ribosome-binding site which lies 5' of the HMW1 gene. A putative transcriptional terminator identical to that identified 3' of the HMW1 ORF is noted, beginning at nucleotide 4804. The discrepancy in the lengths of the two genes is principally accounted for by a 186-bp gap in the HMW2 sequence, beginning at nucleotide position 3839. The derived amino acid sequence of the protein encoded by the HMW2 gene (Figure 4) has a molecular weight of 155,000 and is 71% identical with the derived amino acid sequence of the HMW1 gene.

The derived amino acid sequences of both the HMW1 and HMW2 genes (Figures 2 and 4) demonstrated sequence similarity with the derived amino acid sequence of filamentous hemagglutinin of Bordetella pertussis, a surface-associated protein of this organism. The initial and optimized TFASTA scores for the HMW1-filamentous hemagglutinin sequence comparison were 87 and 186, respectively, with a word size of 2. The z score for the comparison was 45.8. The initial and optimized TFASTA scores for the HMW2-filamentous hemagglutinin sequence comparison were 68 and 196, respectively. The z score for the latter comparison was 48.7. The magnitudes of

the initial and optimized TFASTA scores and the z scores suggested that a biologically significant relationship existed between the HMW1 and HMW2 gene products and filamentous hemagglutinin. When the derived amino acid sequences of HMW1, HMW2, and filamentous hemagglutinin genes were aligned and compared, the similarities were most notable at the amino-terminal ends of the three sequences. Twelve of the first 22 amino acids in the predicted peptide sequences were identical. In addition, the sequences demonstrated a common five-amino-acid stretch, Asn-Pro-Asn-Gly-Ile, and several shorter stretches of sequence identity within the first 200 amino acids.

Example 2:

To further explore the HMW1-filamentous hemagglutinin relationship, the ability of antiserum prepared against the HMW1-4 recombinant protein (rHMW1) to recognize purified filamentous hemagglutinin was assessed. The rHMW1 antiserum demonstrated ELISA reactivity with filamentous hemagglutinin in a dose-dependent manner. Preimmune rabbit serum had minimal reactivity in this assay. The rHMW1 antiserum also was examined in a Western blot assay and demonstrated weak but positive reactivity with purified filamentous hemagglutinin in this system also.

To identify the native Haemophilus protein corresponding to the HMW1 gene product and to determine the extent to which proteins antigenically related to the HMW1 cloned gene product were common among other non-typeable H. influenzae strains, a panel of Haemophilus strains was screened by Western blot with the rHMW1 antiserum. The antiserum recognized both a 125- and a 120-kDa protein band in the homologous strain 12, the putative mature protein products of the HMW1 and HMW2 genes, respectively.

When used to screen heterologous non-typeable H. influenzae strains, rHMW1 antiserum recognized high-molecular-weight proteins in 75% of 125 epidemiologically unrelated strains. In general, the antiserum reacted with one or two protein bands in the 100- to 150-kDa range in each of the heterologous strains in a pattern similar but not identical to that seen in the homologous strain.

Monoclonal antibody X3C is a murine IgG antibody directed against the filamentous hemagglutinin protein of B. pertussis. This antibody can inhibit the binding of B. pertussis cells to Chinese hamster ovary cells and HeLa cells in culture and will inhibit hemagglutination of erythrocytes by purified filamentous hemagglutinin. A Western blot assay was performed in which this monoclonal antibody was screened against the same panel of non-typeable H. influenzae strains discussed above. Monoclonal antibody X3C recognized both the high-molecular-weight proteins in non-typeable H. influenzae strain 12 which were recognized by the recombinant-protein antiserum. In addition, the monoclonal antibody recognized protein bands in a subset of heterologous non-typeable H. influenzae strains which were identical to those recognized by the recombinant-protein antiserum. On occasion, the filamentous hemagglutinin monoclonal antibody appeared to recognize only one of the two bands which had been recognized by the recombinant-protein antiserum. Overall, monoclonal antibody X3C recognized high-molecular-weight protein bands identical to those recognized by the rHMW1 antiserum in approximately 35% of our collection of non-typeable H. influenzae strains.

Example 3:

Mutants deficient in expression of HMW1, MW2 or both proteins were constructed to examine the role of these proteins in bacterial adherence. The following strategy was employed. pHMW1-14 (see Example 1, Figure 5A) was

digested with BamHI and then ligated to a kanamycin cassette isolated on a 1.3-kb BamHI fragment from pUC4K. The resultant plasmid (pHMW1-17) was linearized by digestion with XbaI and transformed into non-typeable H. influenzae strain 12, followed by selection for kanamycin resistant colonies. Southern analysis of a series of these colonies demonstrated two populations of transformants, one with an insertion in the HMW1 structural gene and the other with an insertion in the HMW2 structural gene. One mutant from each of these classes was selected for further studies.

Mutants deficient in expression of both proteins were recovered using the following protocol. After deletion of the 2.1-kb fragment of DNA between two EcoRI sites spanning the 3'-portion of the HMW1 structural gene in pHMW-15, the kanamycin cassette from pUC4K was inserted as a 1.3-kb EcoRI fragment. The resulting plasmid (pHMW1-16) was linearized by digestion with XbaI and transformed into strain 12, followed again by selection for kanamycin resistant colonies. Southern analysis of a representative sampling of these colonies demonstrated that in seven of eight cases, insertion into both the HMW1 and HMW2 loci had occurred. One such mutant was selected for further studies.

To confirm the intended phenotypes, the mutant strains were examined by Western blot analysis with a polyclonal antiserum against recombinant HMW1 protein. The parental strain expressed both the 125-kD HMW1 and the 120-kD HMW2 protein. In contrast, the HMW2⁻ mutant failed to express the 120-kD protein, and the HMW1 mutant failed to express the 125-kD protein. The double mutant lacked expression of either protein. On the basis of whole cell lysates, outer membrane profiles, and colony morphology, the wild type strain and the mutants were otherwise identical with one another. Transmission

electron microscopy demonstrated that none of the four strains expressed pili.

The capacity of wild type strain 12 to adhere to Chang epithelial cells was examined. In such assays, bacteria were inoculated into broth and allowed to grow to a density of $\sim 2 \times 10^9$ cfu/ml. Approximately 2×10^7 cfu were inoculated onto epithelial cell monolayers, and plates were gently centrifuged at $165 \times g$ for 5 minutes to facilitate contact between bacteria and the epithelial surface. After incubation for 30 minutes at 37°C in 5% CO_2 , monolayers were rinsed 5 times with PBS to remove nonadherent organisms and were treated with trypsin-EDTA (0.05% trypsin, 0.5% EDTA) in PBS to release them from the plastic support. Well contents were agitated, and dilutions were plated on solid medium to yield the number of adherent bacteria per monolayer. Percent adherence was calculated by dividing the number of adherent cfu per monolayer by the number of inoculated cfu.

As depicted in Table 1 below (the Tables appear at the end of the descriptive text), this strain adhered quite efficiently, with nearly 90% of the inoculum binding to the monolayer. Adherence by the mutant expressing HMW1 but not HMW2 (HMW2⁻) was also quite efficient and comparable to that by the wild type strain. In contrast, attachment by the strain expressing HMW2 but deficient in expression of HMW1 (HMW1⁻) was decreased about 15-fold relative to the wild type. Adherence by the double mutant (HMW1⁻/HMW2⁻) was decreased even further, approximately 50-fold compared with the wild type and approximately 3-fold compared with the HMW1 mutant. Considered together, these results suggest that both the HMW1 protein and the, HMW2 protein influence attachment to Chang epithelial cells. Interestingly, optimal adherence to this cell line appears to require HMW1 but not HMW2.

Example 4:

Using the plasmids pHMW1-16 and pHMW1-17 (see Example 3) and following a scheme similar to that employed with strain 12 as described in Example 3, three non-typeable Haemophilus strain 5 mutants were isolated, including one with the kanamycin gene inserted into the hmw1-like (designated hmw3) locus, a second with an insertion in the hmw2-like (designated hmw4) locus, and a third with insertions in both loci. As predicted, Western immunoblot analysis demonstrated that the mutant with insertion of the kanamycin cassette into the hmw1-like locus had lost expression of the HMW3 125-kD protein, while the mutant with insertion into the hmw2-like locus failed to express the HMW4 123-kD protein. The mutant with a double insertion was unable to express either of the high molecular weight proteins.

As shown in Table 1 below, wild type strain 5 demonstrated high level adherence, with almost 80% of the inoculum adhering per monolayer. Adherence by the mutant deficient in expression of the HMW2-like protein was also quite high. In contrast, adherence by the mutant unable to express the, HMW1-like protein was reduced about 5-fold relative to the wild type, and attachment by the double mutant was diminished even further (approximately 25-fold). Examination of Giemsa-stained samples confirmed these observations (not shown). Thus, the results with strain 5 corroborate the findings with strain 12 and the HMW1 and HMW2 proteins.

Example 5:

To confirm an adherence function for the HMW1 and HMW2 proteins and to examine the effect of HMW1 and HMW2 independently of other H. influenzae surface structures, the hmw1 and the hmw2 gene clusters were introduced into E. coli DH5 α , using plasmids pHMW1-14 and pHMW2-21, respectively. As a control, the cloning vector, pT7-7, was also transformed into E. coli DH5 α . Western blot

analysis demonstrated that E. coli DH5 α containing the hmw1 genes expressed a 125 kDa protein, while the same strain harboring the hmw2 genes expressed a 120-kDa protein. E. coli DH5 α containing pT7-7 failed to react with antiserum against recombinant HMW1. Transmission electron microscopy revealed no pili or other surface appendages on any of the E. coli strains.

Adherence by the E. coli strains was quantitated and compared with adherence by wild type non-typeable H. influenzae strain 12. As shown in Table 2 below, adherence by E. coli DH5 α containing vector alone was less than 1% of that for strain 12. In contrast, E. coli DH5 α harboring the hmw1 gene cluster demonstrated adherence levels comparable to those for strain 12. Adherence by E. coli DH5 α containing the hmw2 genes was approximately 6-fold lower than attachment by strain 12 but was increased 20-fold over adherence by E. coli DH5 α with pT7-7 alone. These results indicate that the HMW1 and HMW2 proteins are capable of independently mediating attachment to Chang conjunctival cells. These results are consistent with the results with the H. influenzae mutants reported in Examples 3 and 4, providing further evidence that, with Chang epithelial cells, HMW1 is a more efficient adhesin than is HMW2.

Experiments with E. coli HB101 harboring pT7-7, pHMW1-14, or pHMW2-21 confirmed the results obtained with the DH5 α derivatives (see Table 2).

Example 6:

HMW1 and HMW2 were isolated and purified from non-typeable H. influenzae (NTHI) strain 12 in the following manner. Non-typeable Haemophilus bacteria from frozen stock culture were streaked onto a chocolate plate and grown overnight at 37°C in an incubator with 5% CO₂. 50ml starter culture of brain heart infusion (BHI) broth, supplemented with 10 μ g/ml each of hemin and NAD was inoculated with growth on chocolate plate. The start r

culture was grown until the optical density (O.D. - 600nm) reached 0.6 to 0.8 and then the bacteria in the starter culture was used to inoculate six 500 ml flasks of supplemented BHI using 8 to 10 ml per flask. The bacteria were grown in 500 ml flasks for an additional 5 to 6 hours at which time the O.D. was 1.5 or greater. Cultures were centrifuged at 10,000 rpm for 10 minutes.

Bacterial pellets were resuspended in a total volume of 250 ml of an extraction solution comprising 0.5 M NaCl, 0.01 M Na₂EDTA, 0.01 M Tris 50 μ M 1,10-phenanthroline, pH 7.5. The cells were not sonicated or otherwise disrupted. The resuspended cells were allowed to sit on ice at 0°C for 60 minutes. The resuspended cells were centrifuged at 10,000 rpm for 10 minutes at 4°C to remove the majority of intact cells and cellular debris. The supernatant was collected and centrifuged at 100,000 xg for 60 minutes at 4°C. The supernatant again was collected and dialyzed overnight at 4°C against 0.01 M sodium phosphate, pH 6.0.

The sample was centrifuged at 10,000 rpm for 10 minutes at 4°C to remove insoluble debris precipitated from solution during dialysis. The supernatant was applied to a 10 ml CM Sepharose column which has been pre-equilibrated with 0.01 M sodium phosphate, pH 6. Following application to this column, the column was washed with 0.01 M sodium phosphate. Proteins were elevated from the column with a 0 - 0.5M KCl gradient in 0.01 M Na phosphate, pH 6 and fractions were collected for gel examination. Coomassie gels of column fractions were carried out to identify those fractions containing high molecular weight proteins. The fractions containing high molecular weight proteins were pooled and concentrated to a 1 to 3 ml volume in preparation for application of sample to gel filtration column.

A Sepharose CL-4B gel filtration column was equilibrated with phosphate-buffered saline, pH 7.5. The

concentrated high molecular weight protein sample was applied to the gel filtration column and column fractions were collected. Coomassie gels were performed on the column fractions to identify those containing high molecular weight proteins. The column fractions containing high molecular weight proteins were pooled.

The proteins were tested to determine whether they would protect against experimental otitis media caused by the homologous strain.

Chinchillas received three monthly subcutaneous injections with 40 μ g of an HMW1-HMW2 protein mixture in Freund's adjuvant. One month after the last injection, the animals were challenged by intrabullar inoculation with 300 cfu of NTHI strain 12.

Infection developed in 5 of 5 control animals versus 5 of 10 immunized animals. Among infected animals, geometric mean bacterial counts in middle ear fluid 7 days post-challenge were 7.4×10^6 in control animals versus 1.3×10^5 in immunized animals.

Serum antibody titres following immunization were comparable in uninfected and infected animals. However, infection in immunized animals was uniformly associated with the appearance of bacteria down-regulated in expression of the HMW proteins, suggesting bacterial selection in response to immunologic pressure.

Although this data shows that protection following immunization was not complete, this data suggests the HMW adhesin proteins are potentially important protective antigens which may comprise one component of a multi-component NTHI vaccine.

Example 7:

A number of synthetic peptides were derived from HMW1. Antisera then was raised to these peptides. The anti-peptide antisera to peptide HMW1-P5 was shown to recognize HMW1. Peptide HMW1-P5 covers amino acids 1453 to 1481 of HMW1, has the sequence

VDEVIEAKRILEKVKDLSDEEREALAKLG (SEQ ID NO:9), and represents bases 1498 to 1576 in Figure 10.

5 This finding demonstrates that the DNA sequence and the derived protein is being interpreted in the correct reading frame and that peptides derived from the sequence can be produced which will be immunogenic.

SUMMARY OF DISCLOSURE

10 In summary of this disclosure, the present invention provides high molecular weight proteins of non-typeable Haemophilus, genes coding for the same and vaccines incorporating such proteins. Modifications are possible within the scope of this invention.

Table 1. Effect of mutation of high molecular weight proteins on adherence to Chang epithelial cells by nontypable *H. influenzae*.

ADHERENCE*		
Strain	<u>% inoculum</u>	<u>relative to wild type†</u>
Strain 12 derivatives		
wild type	87.7 ± 5.9	100.0 ± 6.7
HMW1- mutant	6.0 ± 0.9	6.8 ± 1.0
HMW2- mutant	89.9 ± 10.8	102.5 ± 12.3
HMW1-/HMW2- mutant	2.0 ± 0.3	2.3 ± 0.3
Strain 5 derivatives		
wild type	78.7 ± 3.2	100.0 ± 4.1
HMW1-like mutant	15.7 ± 2.6	19.9 ± 3.3
HMW2-like mutant	103.7 ± 14.0	131.7 ± 17.8
double mutant	3.5 ± 0.6	4.4 ± 0.8

* Numbers represent mean (\pm standard error of the mean) of measurements in triplicate or quadruplicate from representative experiments.

† Adherence values for strain 12 derivatives are relative to strain 12 wild type; values for strain 5 derivatives are relative to strain 5 wild type.

Table 2. Adherence by *E. coli* DH5 α and HB101 harboring *hmw1* or *hmw2* gene clusters.

Strain*	Adherence relative to <u><i>H. influenzae</i> strain 12[†]</u>
DH5 α (pT7-7)	0.7 \pm 0.02
DH5 α (pHMW1-14)	114.2 \pm 15.9
DH5 α (pHMW2-21)	14.0 \pm 3.7
HB101 (pT7-7)	1.2 \pm 0.5
HB101 (pHMW1-14)	93.6 \pm 15.8
HB101 (pHMW2-21)	3.6 \pm 0.9

* The plasmid pHMW1-14 contains the *hmw1* gene cluster, while pHMW2-21 contains the *hmw2* gene cluster; pT7-7 is the cloning vector used in these constructs.

† Numbers represent the mean (\pm standard error of the mean) of measurements made in triplicate from representative experiments.

CLAIMS

What I claim is:

1. An isolated and purified gene encoding a high molecular weight protein of a non-typeable Haemophilus strain.
2. The gene of claim 1 encoding protein HMW1, HMW2, HMW3 or HMW4 or a variant or fragment of said protein retaining the immunological ability to protect against disease caused by a non-typeable Haemophilus strain.
3. The gene of claim 2 having the DNA sequence shown in Figure 1 and encoding protein HMW1 having the derived amino acid sequence of Figure 2.
4. The gene of claim 2 having the DNA sequence shown in Figure 3 and encoding protein HMW2 having the derived amino acid sequence of Figure 4.
5. The gene claimed in claim 2 having the partial DNA sequence shown in Figure 8 and encoding protein HMW3 having the derived amino acid sequence of Figure 10.
6. The gene claimed in claim 2 having the partial DNA sequence shown in Figure 9 and encoding protein HMW4 having the derived amino acid sequence of Figure 10.
7. A purified and isolated gene cluster comprising a nucleotide sequence for a structural gene encoding a high molecular weight protein of a non-typeable Haemophilus strain and at least one downstream nucleotide sequence for an accessory gene for effecting expression of a gene product fully encoded by said structural gene.
8. The gene cluster claimed in claim 7 comprising a DNA sequence coding for protein HMW1 or HMW2 and two downstream accessory genes.
9. The gene cluster of claim 8 having the DNA sequence shown in Figure 6.
10. The gene cluster of claim 8 having the DNA sequence shown in Figure 7.
11. A high molecular weight protein of non-typeable Haemophilus which is encoded by a gene as defined in

claim 1, or any variant or fragment thereof retaining the immunological ability to protect against disease caused by a non-typeable Haemophilus strain.

12. The protein of claim 11 which is HMW1 encoded by the DNA sequence shown in Figure 1, having the derived amino acid sequence of Figure 2 and having an apparent molecular weight of 125 kDa.

13. The protein claim 11 which is HMW2 encoded by the DNA sequence shown in Figure 3 and having the derived amino acid sequence of Figure 4 and having an apparent molecular weight of 120 kDa.

14. An isolated and purified high molecular weight protein of non-typeable Haemophilus influenzae which is antigenically related to the filamentous hemagglutinin surface protein of Bordetella pertussis.

15. The protein of claim 14 which is HMW1, HMW2, HMW3 or HMW4.

16. A conjugate comprising a protein as claimed in claim 11 or 14 linked to a antigen, hapten or polysaccharide for eliciting an immune response to said antigen, hapten or polysaccharide.

17. The conjugate as claimed in claim 16 wherein said polysaccharide is a protective polysaccharide against Haemophilus influenzae type b.

18. A synthetic peptide having an amino acid sequence corresponding to at least one protective epitope of a high molecular weight protein of non-typeable Haemophilus influenzae.

19. The peptide of claim 18 wherein said protein is HMW1, HMW2, HMW3 or HMW4.

FIG. 1A. DNA SEQUENCE OF HIGH MOLECULAR WEIGHT PROTEIN

I (HMW1)

1 ACAGCGTTCT CTTAATACTA GTACAAACCC ACAATAAAT ATGACAAACA
 51 ACAATTACAA CACCTTTTTT GCAGTCTATA TGCAAAATATT TTAATAAATA
 101 GTATAAATCC GCCATATAAA ATGGTATAAT CTTTCATCTT TCATCTTTCA
 151 TCTTTTCATCT TTCATCTTTC ATCTTTTCATC TTTTCATCTTT CATCTTTTCAT
 201 CTTTTCATCTT TCATCTTTCA TCTTTTCATCT TTCATCTTTC ACATGCCCTG
 251 ATGAACCGAG GGAAGGAGG GAGGGCAAG AATGAAGAGG GAGCTGAACG
 301 AACGCAAAATG ATAAAGTAAT TTAATTGTTC AACTAACCTT AGGAGAAAAT
 351 ATGAACAAGC TATATCGTCT CAAATTCAGC AAACGCCCTGA ATGCTTTGGT
 401 TGCTGTGTCT GAATTGGCAC GGGGTTGTGA CCATTCCACA GAAAAAGGCA
 451 GCGAAAAACC TGCTCGCATG AAAGTGGCTC ACTTAGCGTT AAAGCCACTT
 501 TCCGCTATGT TACTATCTTT AGGTGTAACA TCTATTCCAC AATCTGTTTT
 551 AGCAAGCGGC TTACAAGGAA TGGATGTAGT ACACGGCACA GCCACTATGC
 601 AAGTAGATGG TAATAAAACC ATTATCCGCA ACAGTGTGA CGATATCATT
 651 AATTGGAAAC AATTAAACAT CGACCAAAAT GAAATGGTGC AGTTTTTACA
 701 AGAAAACAAC AACTCCGCCG TATTCAACCG TGTTACATCT AACCAAATCT

1 / 88

FIG. 1B.

751 CCCAATTAAA AGGGATTTTA GATTCTAACG GACAAGTCTT TTTAATCAAC
 801 CCAAAATGGTA TCACAATAGG TAAAGACGCA ATTATTAAACA CTAATGGCTT
 851 TACGGGCTTCT ACGCTAGACA TTTCTAACGA AAACATCAAG GCGCGTAATT
 901 TCACCC TTCGA GCAAACCAA GATAAAGCGC TCGCTGAAAT TGTGAATCAC
 951 GGTTTAATTA CTGTCGGTAA AGACGGCAGT GTAAATCTTA TTGGTGGCAA
 1001 AGTGAAAAAC GAGGGTGTGA TTAGCGTAAA TGGTGGCAGC ATTTCTTTAC
 1051 TCGCAGGGCA AAAAATCACC ATCAGCGATA TAATAAACCC AACCATTA
 1101 TACAGCATTG CCGCGCCTGA AAATGAAGCG GTCAATCTGG GCGATATTTT
 1151 TGCCAAAGGC GGTAACATTA ATGTCCGTGC TGCCACTATT CGAAACCAAG
 1201 GTAAACTTTC TGCTGATTCT GTAAGCAAAG ATAAAAGCGG CAATATTGTT
 1251 CTTTCCGCCA AAGAGGGTGA AGCGGAAATT GCGGGTGTA TTTCCGCTCA
 1301 AAATCAGCAA GCTAAAGCG GCAAGCTGAT GATTACAGGC GATAAAGTCA
 1351 CATTAAAAAC AGGTGCAGTT ATCGACCTTT CAGGTAAAGA AGGGGGAGAA
 1401 ACTTACCCTTG GCGGTGACGA GCGCGGCGAA GGTA AAAAGG GCATTCAATT
 1451 AGCAAAGAAA ACCTCTTTAG AAAAAGGCTC AACCATCAAT GTATCAGGCA
 1501 AAGAAAAAGG CGGACCGGCT ATTGTGTGG GCGATATTGC GTTAATTGAC

FIG. 1C.

1551	GGCAATATTA	ACGCTCAAGG	TAGTGGTGAT	ATCGCTAAAA	CCGGTGGTTT
1601	TGTGGAGACG	TCGGGGCATG	ATTTATTTCAT	CAAAGACAAAT	GCAATTGTTG
1651	ACGCCAAAGA	GTGGTTGTTA	GACCCGGATA	ATGTATCTAT	TAATGCAGAA
1701	ACAGCAGGAC	GCAGCAATAC	TTCAGAAGAC	GATGAATACA	CGGGATCCGG
1751	GAATAGTGCC	AGCACCCCAA	AACGAAACAA	AGAAAAGACA	ACATTAAACAA
1801	ACACAACTCT	TGAGAGTATA	CTAAAAAAG	GTACCTTTGT	TAACATCACT
1851	GCTAATCAAC	GCATCTATGT	CAATAGCTCC	ATTAATTTAT	CCAATGGCAG
1901	CTTAACTCTT	TGGAGTGAGG	GTCGGAGCGG	TGGCGGCGTT	GAGATTAAACA
1951	ACGATATTAC	CACCGGTGAT	GATACCAGAG	GTGCAAACTT	AACAAATTTAC
2001	TCAGGGGGCT	GGGTGATGT	TCATAAAAAT	ATCTCACTCG	GGGCGCAAGG
2051	TAACATAAAC	ATTACAGCTA	AACAAGATAT	CGCCTTTGAG	AAAGGAAGCA
2101	ACCAAGTCAT	TACAGGTCAA	GGGACTATTA	CCTCAGGCAA	TCAAAAAGGT
2151	TTTAGATTTA	ATAATGTCTC	TCTAAACGGC	ACTGGCAGCG	GACTGCAATT
2201	CACCACTAAA	AGAACCAATA	AATACGCTAT	CACAAATAAA	TTTGAAGGGA
2251	CTTTAAATAT	TTCAGGGGAA	GTGAACATCT	CAATGGTTTT	ACCTAAAAAT
2301	GAAAGTGGAT	ATGATAAATT	CAAAGGACGC	ACTTACTGGA	ATTTAACCTC

3 / 8

FIG. 1D.

2351	CTTAAATGTT	TCCGAGAGTG	GCGAGTTTAA	CCTCACTATT	GACTCCAGAG
2401	GAAGCGATAG	TGCAGGCACA	CTTACCCAGC	CTTATAAATT	AAACGGTATA
2451	TCATTCAACA	AAGACACTAC	CTTTAATGTT	GAACGAAATG	CAAGAGTCAA
2501	CTTTGACATC	AAGGCACCAA	TAGGGATAAA	TAAGTATTCT	AGTTTGAAAT
2551	ACGCATCATT	TAATGGAAAC	ATTTCAGTTT	CGGGAGGGGG	GAGTGTGTGAT
2601	TTACACACTTC	TCGCCCTCATC	CTCTAACGTC	CAAACCCCGG	GTGTAGTTAT
2651	AAATTCTAAA	TACTTTAATG	TTTCAACACAGG	GTCAAGTTTA	AGATTTAAAA
2701	CTTCAGGCTC	AACAAAAACT	GGCTTCTCAA	TAGAGAAAGA	TTTAACTTTA
2751	AATGCCACCG	GAGGCAACAT	AACACTTTTG	CAAGTTGAAG	GCACCGATGG
2801	AATGATTGGT	AAAGGCATTG	TAGCCAAAAA	AAACATAACC	TTTGAAGGAG
2851	GTAACATCAC	CTTTGGCTCC	AGGAAAGCCG	TAACAGAAAT	CGAAGGCAAT
2901	GTTACTATCA	ATAACAACGC	TAAAGTCACT	CTTATCGGTT	CGGATTTTGA
2951	CAACCATCAA	AAACCTTTAA	CTATTAAAAA	AGATGTCATC	ATTAATAGCG
3001	GCAACCTTAC	CGCTGGAGGC	AATATTGTCA	ATATAGCCGG	AAATCTTACC
3051	GTTGAAAGTA	ACGCTAATT	CAAAGCTATC	ACAAAATTCA	CTTTTAATGT
3101	AGGCGGCTTG	TTTGACAACA	AAGGCAATC	AAATATTTC	ATTGCCAAAG
3151	GAGGGGCTCG	CTTTAAAGAC	ATTGATAATT	CCAAGAAATT	AAGCATCACC

FIG. 1E.

3201	ACCAACTCCA	GCTCCACTTA	CCGCACTATT	ATAAGCGGCA	ATATAACCAA
3251	TAAAAACGGT	GATTTAAATA	TTACGAACGA	AGGTAGTGAT	ACTGAAATGC
3301	AAATTGGCGG	CGATGTCTCG	CAAAAAGAAG	GTAATCTCAC	GATTTCTTCT
3351	GACAAAATCA	ATATTACCAA	ACAGATAACA	ATCAAGGCAG	GTGTTGATGG
3401	GGAGAAATTC	GATTCAGACG	CGACAAACAA	TGCCAATCTA	ACCATTAAAA
3451	CCAAGAATTT	GAAATTAAACG	CAAGACCTAA	ATATTTCAGG	TTTCAATAAA
3501	GCAGAGATTA	CAGCTAAAGA	TGGTAGTGAT	TTAACTATTG	GTAACACCAA
3551	TAGTGCTGAT	GGTACTAATG	CCAAAAAAGT	AACCTTTAAC	CAGGTTAAAG
3601	ATTCAAAAAT	CTCTGCTGAC	GGTCACAAGG	TGACACTACA	CAGCAAAGTG
3651	GAAACATCCG	GTAGTAATAA	CAACACTGAA	GATAGCAGTG	ACAATAATGC
3701	CGGCTTAACT	ATCGATGCAA	AAAATGTAAC	AGTAAACAAC	AATATTACTT
3751	CTCACAAAGC	AGTGAGCATC	TCTGCGACAA	GTGGAGAAAT	TACCACATAAA
3801	ACAGGTACAA	CCATTAAACG	AACCACTGGT	AACGTGGAGA	TAACCGCTCA
3851	AACAGGTAGT	ATCCTAGGTG	GAAATTGAGTC	CAGCTCTGGC	TCTGTAACAC
3901	TTACTGCAAC	CGAGGGCGCT	CTTGCTGTAA	GCAATATTTC	GGGCAACACC
3951	GTTACTGTTA	CTGCAAATAG	CGGTGCATTA	ACCACTTTGG	CAGGCTCTAC

FIG. 1F.

4001 AATTAAAGGA ACCGAGAGTG TAACCACTTC AAGTCAATCA GCGGATATCG

4051 GCGGTACGAT TTCTGGTGGC ACAGTAGAGG TTAAAGCAAC CGAAAGTTTA

4101 ACCACTCAAT CCAATTCAAA AATTAAAGCA ACAACAGGCG AGGCTAACGT

4151 AACAAAGTGA ACAGGTACAA TTGGTGGTAC GATTTCGGT AATACGGTAA

4201 ATGTTACGGC AAACGCTGGC GATTTAACAG TTGGGAATGG CGCAGAAATT

4251 AATGCGACAG AAGGAGCTGC AACCTTAACT ACATCATCGG GCAAATTAAC

4301 TACCGAAGCT AGTTCACACA TTACTIONCAG CAAGGGTCAG GTAAATCTTT

4351 CAGCTCAGGA TGGTAGCGTT GCAGGAAGTA TTAATGCCGC CAATGTGACA

4401 CTAAATACTA CAGGCACTTT AACTACCGTG AAGGGTTCAA ACATTAATGC

4451 AACCAGCGGT ACCTTGGTTA TTAACGCAAA AGACGCTGAG CTAAATGGCG

4501 CAGCATTTGG TAACCACACA GTGGTAAATG CAACCAACGC AAATGGCTCC

4551 GGCAGCGTAA TCGCGACAAC CTCAAGCAGA GTGAACATCA CTGGGGATTT

4601 AATCACAAATA AATGGATTAA ATATCATTTT AAAAAACGGT ATAAACACCG

4651 TACTGTATAA AGGCGTTAAA ATTGATGTGA AATACATTCA ACCGGGTATA

4701 GCAAGCGTAG ATGAAGTAAT TGAAGCGAAA CGCATCCTTG AGAAGGTAAA

4751 AGATTTATCT GATGAAGAAA GAGAAGCGTT AGCTAAACTT GGAGTAAAGTG

4801 CTGTACGTTT TATTGAGCCA AATAATACAA TTACAGTCGA TACACAAAAT

6/68

7/68

FIG. 1G.

4851 GAATTGCAA CCAGACCAT T AAGTCGAATA GTGATTTC TG AAGGCAGGGC
4901 GTGTTTCTCA AACAGTGATG GCGCGACGGT GTGCCGTTAAT ATCGCTGATA
4951 ACGGGCGGTA GCGGTCAGTA ATTGACAAGG TAGATTTCAT CCTGCAATGA
5001 AGTCATTTTA TTTTCGTATT ATTTACTGTG TGGGTTAAAG TTCAGTACGG
5051 GCTTTACCCA TCTTGTA AAA AATTACGGAG AATACAATAA AGTATTTTA
5101 ACAGGTTATT ATTATG

FIG. 2A. AMINO ACID SEQUENCE OF HIGH MOLECULAR WEIGHT

PROTEIN I

1 MNKIYRLKFS KRLNALVAVS ELARGCDHST EKGSEKPARM KVRHLALKPL
51 SAML LSLGVT SIPQSVLASG LQMDVVHGT ATMQVDGNKT IIRNSVDAIL
101 NWKQFNIDQN EMVQFLQENN NSAVFNRVTS NQISQLKGIL DSNQGVFLIN
151 PNGITIGKDA IINTNGFTAS TLDISNENIK ARNFTFEQTK DKALAEIVNH
201 GLITTVGKDG VNLIGGKVKV EGVISVNGGS ISLLAGQKIT ISDIINPTIT
251 YSIAAPENEA VNLGDIFAKG GNINVRAATI RNQKLSADS VSKDKSGNIV
301 LSAKEGEAEI GGVIS AQNQQ AKGKLMITG DKVTLKTGAV IDLSGKEGGE
351 TYLGGDERGE GKNGIQ LAKK TSLEKGSTIN VSGKEKGRA IVWGDIALID
401 GNINAQGS D IAKTGGFVET SGHDLFIKDN AIVDAKEWLL DFDNVSINAE
451 TAGRSNTSED DEYTGSGNSA STPKRNKEKT TLTNTTLESI LKKGTFVNIT
501 ANQRIYVNSS INLSNGSLTL WSEGRSGGV EINNDITTD DTRGANLTIY
551 SGGWVDVHKN ISLGAQGNIN ITAKQDIAFE KGSNQVITGQ GTITSGNQKG
601 FRFNNVSLNG TGSGLQFTTK RTNKYAITNK FEGTLNISGK VNISMVLPKN
651 ESGYDKFKGR TYWNLTSLNV SESGEFNLT I DSRGSDSAGT LTQPYNLNGI
701 SFNKDTTFNV ERNARVNFDI KAPIGINKYS SLNYASFNGN ISVSGGGSVD

FIG. 2B.

751	FTLLASSNV	QTPGVVINSK	YFNVSTGSSL	RFKTSGSTKT	GFSIEKDLTL
801	NATGGNITLL	QVEGTDGMIG	KGIVAKKNIT	FEGGNITFGS	RKAVTEIEGN
851	VTINNANVT	LIGSDFDNHQ	KPLTIKKDVI	INSGNLTAGG	NIVNIAGNLT
901	VESNANFKAI	TNFTFNVGGL	FDNKGNSNIS	IAKGARFKD	IDNSKNLSIT
951	TNSSSTYRTI	ISGNITNKNG	DLNITNEGSD	TEMQIGGDVS	QKEGNLTISS
1001	DKINITKQIT	IKAGVDGENS	DSDATNNANL	TIKTKEKLT	QDLNISGFNK
1051	AEITAKDGSD	LTIGNTNSAD	GTNAKKVTFN	QVKDSKISAD	GHKVTLHSKV
1101	ETSGSNNNTE	DSSDNNAGLT	IDAKNVTVNN	NITSHKAVSI	SATSGEITTK
1151	TGTTINATTG	NVEITAQTS	ILGGIESSG	SVTLTATEGA	LAVSNISGNT
1201	VTVTANS GAL	TTLAGSTIKG	TESVTTSSQS	GDIGGTISGG	TVEVKATESL
1251	TTQSNSKIIKA	TTGEANVTSA	TGTIGGTISG	NTVNVVTANAG	DLTVGNGAEI
1301	NATEGAATLT	TSSGKLTEA	SSHITSAKGQ	VNLSAQDGSV	AGSINAANVT
1351	LNTTGTLTTV	KGSNINATSG	TLVINAKDAE	LNGAALGNHT	VVNATNANGS
1401	GSVIATTSSR	VNITGDLITI	NGLNIISKNG	INTVLLKGVK	IDVKYIQPGI
1451	ASVDEVIEAK	RILEKVKDLS	DEEREALAKL	GVSAVRFIEP	NNTITVDTQN
1501	EFATRPLSRI	VISEGRACFS	NSDGATVCVN	IADNGR	

9/08

FIG. 3A. AMINO ACID SEQUENCE OF HIGH MOLECULAR WEIGHT

PROTEIN II (HMW2)

1 TAAATATACA AGATAATAAA AATAAATCAA GATTTTGTG ATGACAAACA
 51 ACAATTACAA CACCTTTTTC GCAGTCTATA TGCAAATATT TTAAAAAAT
 101 AGTATAAATC CGCCATATAA AATGGTATAA TCCTTTCATCT TTCATCTTTA
 151 ATCTTTCATC TTTTCATCTTT CATCTTTCAT CTTTCATCTT TCATCTTTCA
 201 TCTTTTCATCT TTCATCTTTC ATCTTTCATC TTTTCATCTTT CACATGAAAT
 251 GATGAACCGA GGAAGGGAG GGAGGGCAA GAATGAAGAG GGAGCTGAAC
 301 GAACGCAAAAT GATAAAGTAA TTTAATTGTT CAACTAACCT TAGGAGAAA
 351 TATGAACAAG ATATATCGTC TCAAATTTCAG CAAACGCCCTG AATGCTTTGG
 401 TTGCTGTGTC TGAATTGGCA CGGGTTGTG ACCATTCCAC AGAAAAAGGC
 451 TTCCGCTATG TTAATATCTT TAGGTGTAAC CACTTAGCGT TAAAGCCACT
 501 TTCCGCTATG TTAATATCTT TAGGTGTAAC ATCTATTCCA CAATCTGTTT
 551 TAGCAAGCGG CTTACAAGGA ATGGATGTAG TACACGGCAC AGCCACTATG
 601 CAAGTAGATG GTAATAAAAC CATTATCCGC AACAGTGTG ACGCTATCAT
 651 TAATTGGAAA CAATTTAACA TCGACCAAAA TGAATGGTG CAGTTTTTAC
 701 AAGAAAACAA CAACTCCGCC GTATTCAACC GTGTTACATC TAACCAATC

FIG. 3B.

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751  TCCCAATTAA AAGGATTTT AGATTCTAAC GGACAAGTCT TTTTAATCAA
801  CCCAAATGGT ATCACAATAG GTAAAGACGC AATTATTAACT ACTAATGGCT
851  TTACGGCTTC TACGCTAGAC ATTTCTAACG AAAACATCAA GCGCGTAAT
901  TTCACCTTCG AGCAAACCAA AGATAAAGCG CTCGCTGAAA TTGTGAATCA
951  CGGTTTAATT ACTGTCGGTA AAGACGGCAG TGTAATCTT ATTGGTGGCA
1001 AAGTGAAAAA CGAGGGGTG ATTAGCGTAA ATGGTGGCAG CATTCTTTA
1051 CTCGCAGGC AAAAAATCAC CATCAGCGAT ATAATAAACC CAACCATTA
1101 TTACAGCATT GCCGCGCCTG AAAATGAAGC GGTCATCTG GCGATATT
1151 TTGCCAAAGG CGGTAACATT AATGTCCGTG CTGCCACTAT TCGAAAACCAA
1201 GGTAACCTT CTGCTGATTC TGTAAGCAA GATAAAGCG GCAATATTGT
1251 TCTTTCGCC AAAGAGGGTG AAGCGGAAAT TGGCGGTGTA ATTTCCGCTC
1301 AAAATCAGCA AGCTAAAGGC GGCAAGCTGA TGATTACAGG CGATAAAGTC
1351 ACATTAAAAA CAGGTGCAGT TATCGACCTT TCAGGTAAAG AAGGGGAGA
1401 AACTTACCTT GCGGTGACG AGCGCGCGCA AGGTAAAAAC GGCATTCAAT
1451 TAGCAAAGAA AACCTCTTTA GAAAAAGGCT CAACCATCAA TGTATCAGGC
1501 AAAGAAAAAG GCGACGCGC TATTGTGTG GCGGATATTG CGTTAATTGA

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FIG. 3C.

1551	CGGCAATATT AACGCTCAAG GTAGTGGTGA TATCGCTAAA ACCGGTGGTT	12 / 68
1601	TTGTGGAGAC ATCGGGGCAT TATTTATCCA TTGACAGCAA TGCAATTGTT	
1651	AAAACAAAAG AGTGGTTGCT AGACCCCTGAT GATGTAACAA TTGAAGCCGA	
1701	AGACCCCCCTT CGCAATAATA CCGGTATAAA TGATGAATTC CCAACAGGCA	
1751	CCGGTGAAGC AAGCGACCCT AAAAAAATA GCGAACTCAA AACACGCTA	
1801	ACCAATACAA CTATTTCAAATTATCTGAAA AACGCCCTGGA CAATGAATAT	
1851	AACGGCATCA AGAAAACTTA CCGTTAATAG CTCAATCAAC ATCGGAAGCA	
1901	ACTCCCACCTT AATTCTCCAT AGTAAAGGTC AGCGTGGCGG AGGCGTTCAG	
1951	ATTGATGGAG ATATTACTTC TAAAGGCGGA AATTTAACCA TTTATTCTGG	
2001	CGGATGGGTT GATGTTCATA AAAATATTAC GCTTGATCAG GGTTTTTTAA	
2051	ATATTACCGC CGCTTCCGTA GCTTTTGAAG GTGGAATAA CAAAGCACGC	
2101	GACGCGGCAA ATGCTAAAAAT TGTGCCCCAG GGCACGTGTA CCATTACAGG	
2151	AGAGGGGAAA GATTTCAGGG CTAACAACGT ATCTTTAAAC GGAACGGGTA	
2201	AAGGTCTGAA TATCATTTCA TCAGTGAATA ATTTAACCCA CAATCTTAGT	
2251	GGCACAAATTA ACATATCTGG GAATATAACA ATTAACCCAA CTACGAGAAA	
2301	GAACACCTCG TATTGGCAA CCAGCCATGA TTCGCACTGG AACGTCAGTG	
2351	CTCTTAATCT AGAGACAGGC GCAAATTTTA CCTTTATTAA ATACATTCA	

FIG. 3D.

2401	AGCAATAGCA	AAGGCTTAAC	AACACAGTAT	AGAAGCTCTG	CAGGGGTGAA
2451	TTTTAACGGC	GTAAATGGCA	ACATGTCATT	CAATCTCAA	GAAGGAGCGA
2501	AAGTTAATTT	CAAATTAAAA	CCAAACGAGA	ACATGAACAC	AAGCAAACCT
2551	TTACCAATTC	GGTTTTTAGC	CAATATCACA	GCCACTGGTG	GGGGCTCTGT
2601	TTTTTTTGAT	ATATATGCCA	ACCATTCCTG	CAGAGGGGCT	GAGTTAAAAA
2651	TGAGTGAAAT	TAATATCTCT	AACGGCGCTA	ATTTTACCTT	AAATTCCCCT
2701	GTTCGCGGCG	ATGACGCTTT	TAAAATCAAC	AAAGACTTAA	CCATAAATGC
2751	AACCAATTCA	AATTCAGCC	TCAGACAGAC	GAAAGATGAT	TTTTATGACG
2801	GGTACGCACG	CAATGCCATC	AATTCAACCT	ACAACATATC	CATTCTGGGC
2851	GGTAATGTCA	CCCTTGGTGG	ACAAAACCTCA	AGCAGCAGCA	TTACGGGGAA
2901	TATTACTATC	GAGAAAGCAG	CAAATGTTAC	GCTAGAAGCC	ATAACGCCCC
2951	CTAATCAGCA	AAACATAAGG	GATAGAGTTA	TAAAACCTGG	CAGCTTGCTC
3001	GTTAATGGGA	GTTTAAAGTTT	AACTGGCGAA	AATGCAGATA	TTAAAGGCAA
3051	TCTCACTATT	TCAGAAAGCG	CCACTTTTAA	AGGAAAGACT	AGAGATACCC
3101	TAAATATCAC	CGGCAATTTT	ACCAATAATG	GCACTGCCGA	AATTAATATA
3151	ACACAAGGAG	TGGTAAAAC	TGGCAATGTT	ACCAATGATG	GTGATTTAAA

FIG. 3E.

3201 CATTACCACT CACGCTAAAC GCAACCAAAG AAGCATCATC GCGGAGATA
3251 TAATCAACAA AAAAGGAAGC TTAATATATTA CAGACAGTAA TAATGATGCT
3301 GAAATCCAAA TTGGCGGCAA TATCTCGCAA AAAGAAGGCA ACCTCACGAT
3351 TTCTTCCGAT AAAATTAAATA TCACCAAACA GATAACAATC AAAAAGGGTA
3401 TTGATGGAGA GGACTCTAGT TCAGATGCGA CAAGTAATGC CAACCTAACT
3451 ATTAAAACCA AAGAATTGAA ATTGACAGAA GACCTAAGTA TTTCAGGTTT
3501 CAATAAAGCA GAGATTACAG CCAAAGATGG TAGAGATTTA ACTATTGGCA
3551 ACAGTAATGA CGGTAACAGC GGTGCCGAAG CCAAAACAGT AACTTTTAAC
3601 AATGTTAAAG ATTCAAAAAT CTCTGCTGAC GGTCACAATG TGACACTAAA
3651 TAGCAAAGTG AAAACATCTA GCAGCAATGG CGGACGTGAA AGCAATAGCG
3701 ACAACGATAC CGGCTTAACT ATTACTGCAA AAAATGTAGA AGTAAACAAA
3751 GATATTACTT CTCTCAAAAC AGTAAATATC ACCGCGTCGG AAAAGGTTAC
3801 CACCACAGCA GGCTCGACCA TTAACGCAAC AAATGGCAA GCAAGTATTA
3851 CAACCAAAAC AGGTGATATC AGCGGTACGA TTTCCGGTAA CACGGTAAGT
3901 GTTAGCGCGA CTGGTGATTT AACCACATAA TCCGGCTCAA AAATTGAAGC
3951 GAAATCGGGT GAGGCTAATG TAACAAGTGC AACAGGTACA ATTGGCCGTA

14 / 68

FIG. 3F.

4001 CAATTTCCGG TAATACGGTA AATGTTACGG CAAACGCTGG CGATTTAACA
 4051 GTTGGGAATG GGCAGAAAT TAATGCGACA GAAGGAGCTG CAACCTTAAC
 4101 CGCAACAGGG AATACCTTGA CTA CTGAAGC CGGTTCTAGC ATCACTTCAA
 4151 CTAAGGGTCA GGTAGACCTC TTGGCTCAGA ATG GTAGCAT CGCAGGAAGC
 4201 ATTAATGCTG CTAATGTGAC ATTAAATACT ACAGGCACCT TAACCACCGT
 4251 GGCAGGCTCG GATATTAAAG CAACCAGCGG CACCTTGCTT ATTAACGCAA
 4301 AAGATGCTAA GCTAAATGGT GATGCATCAG GTGATAGTAC AGAAGTGAAT
 4351 GCAGTCAACG CAAGCGGCTC TGGTAGTGTG ACTGCGGCAA CCTCAAGCAG
 4401 TGTGAATATC ACTGGGGATT TAAACACAGT AAATGGGTTA AATATCATTT
 4451 CGAAAGATGG TAGAAACACT GTGCGCTTAA GAGCAAGGA AATTGAGGTG
 4501 AAATATATCC AGCCAGGTGT AGCAAGTGTA GAAGAAGTAA TTGAAGCGAA
 4551 ACGCGTCCTT GAAAAGTAA AAGATTATC TGATGAAGAA AGAGAAACAT
 4601 TAGCTAAACT TGGTGTAAGT GCTGTACGTT TTGTTGAGCC AAATAATACA
 4651 ATTACAGTCA ATACACAAA TGAATTTACA ACCAGACCGT CAAGTCAAGT
 4701 GATAATTTCT GAAGGTAAGG CGTGTTTCTC AAGTGTAAT GGCGACGAG
 4751 TATGTACCAA TGTGCTGAC GATGGACAGC CGTAGTCAGT AATTGACAAG
 4801 GTAGATTTCA TCCTGCAATG AAGTCATTTT ATTTTCGTAT TATTACTGT

15 / 60

16/68

FIG. 3G.

4851 GTGGGTAAA GTTCAGTACG GGCTTACCC ATCTTGTAAG AAATTACGGA
4901 GAATACAATA AAGTATTTT AACAGGTAT TATTATG

FIG. 4A. AMINO ACID SEQUENCE OF HIGH MOLECULAR WEIGHT

PROTEIN 2

1 MNKIYRLKFS KRLNALVAVS ELARGCDHST EKGSEKPARM KVRHLALKPL
51 SAML LSLGVT SIPQSVLASG LQGM DVVHGT ATMQVDGNKT IIRNSVDAIL
101 NWKQFNIDQN EMVQFLQENN NSAVFN RVTS NQISQLKGIL DSNQVFLIN
151 PNGITIGKDA IINTNGFTAS TLDISNENIK ARNFTFEQTK DKALAEIVNH
201 GLITVGKDG S VNLIGGKVKN EGVISVNGGS ISLLAGQKIT ISDIINPTIT
251 YSIAAPENEA VNLGDIFAKG GNINVRAATI RNQKLSADS VSKDKSGNIV
301 LSAKEGEAEI GGVIS AQNQQ AKGKLMITG DKVTLKTGAV IDLSGKEGGE
351 TYLGGDERGE GKNIGIQLAKK TSLEKGSTIN VSGKEKGGRA IVWGDIALID
401 GNINAQGS GD IAKTGGFVET SGHDLFIKDN AIVDAKEWLL DFDNV SINAE
451 DPLRNNTGIN DEFPTGTGEA SDPKKNSELK TTLTNTTISN YLKNAWTMNI
501 TASRKLT VNS SINIGSNSHL ILHSGQRGG GVQIDGDITS KGNLTIYSG
551 GWVDVHK NIT LDQGF LNITA ASVAFEGGNN KARDAANAKI VAQGTVTITG
601 EGKDFRANNV SLNGTGKGLN IISVNNLTH NLSGTINISG NITINQTRK
651 NTSYWQTS HD SHWNVSALNL ETGANFTFIK YISSNSKGLT TQYRSSAGVN
701 FNGVNGNMSF NLKEGAKVNF KLKPENMMNT SKPLPIRFLA NITATGGGSV

17/68

FIG. 4B.

751 FFDIYANHSG RGAELKMSEI NISNGANFTL NSHVRGDDAF KINKDLTINA
 801 TNSNFSLRQT KDDFYDGYAR NAINSTYNIS ILGGNVTLGG QNSSSSITGN
 851 ITIEKAANVT LEANNAPNQQ NIRDVRIKLG SLLVNGSLSL TGENADIKGN
 901 LTISESATFK GKTRDTLNIIT GNFTNNGTAE INITQGVVKL GNVNDGDGLN
 951 ITHAKRNQR SIIIGDIIINK KGSLNITDSN NDAEIQIGGN ISQKEGNLTI
 1001 SSDKINITKQ ITIKKGIDGE DSSSDATSNALTLTKTELKLTEDLSISGF
 1051 NKAELITAKDG RDLTIGNSND GNSGAEAKTVTFNNVKDSKISADGHNVTLN
 1101 SKVKTSSSNG GRESNSDNDTGLTITAKNVEVNKDITSLKTVNITASEKVT
 1151 TTAGSTINATNGKASITTKTGDISGTISGNTVSVSATVDLTTKSGSKIEA
 1201 KSGEANVTSA TGTIGGTISGNTVNVNANAGDLTVGNAGAEINATEGAATLT
 1251 ATGNTLTTEAGSSITSTKGQVDLLAQNGSIAGSINAANVTLNTTGTLTTV
 1301 AGSDIKATSGTLVINAKDAKLNGDASGDSTEVNAVNASGSGSVTAATSSS
 1351 VNITGDLNTVNGLNIISKDGRNTVRLRGKEIEVKYIQPGVASVEEVIEAK
 1401 RVLEKVKDLSDEERETLAKLGVSAVRFVEPNNTITVNTQNFFTTRPSSQV
 1451 IISEGKACFSSGNGARVCTNVADDGQP

19/68

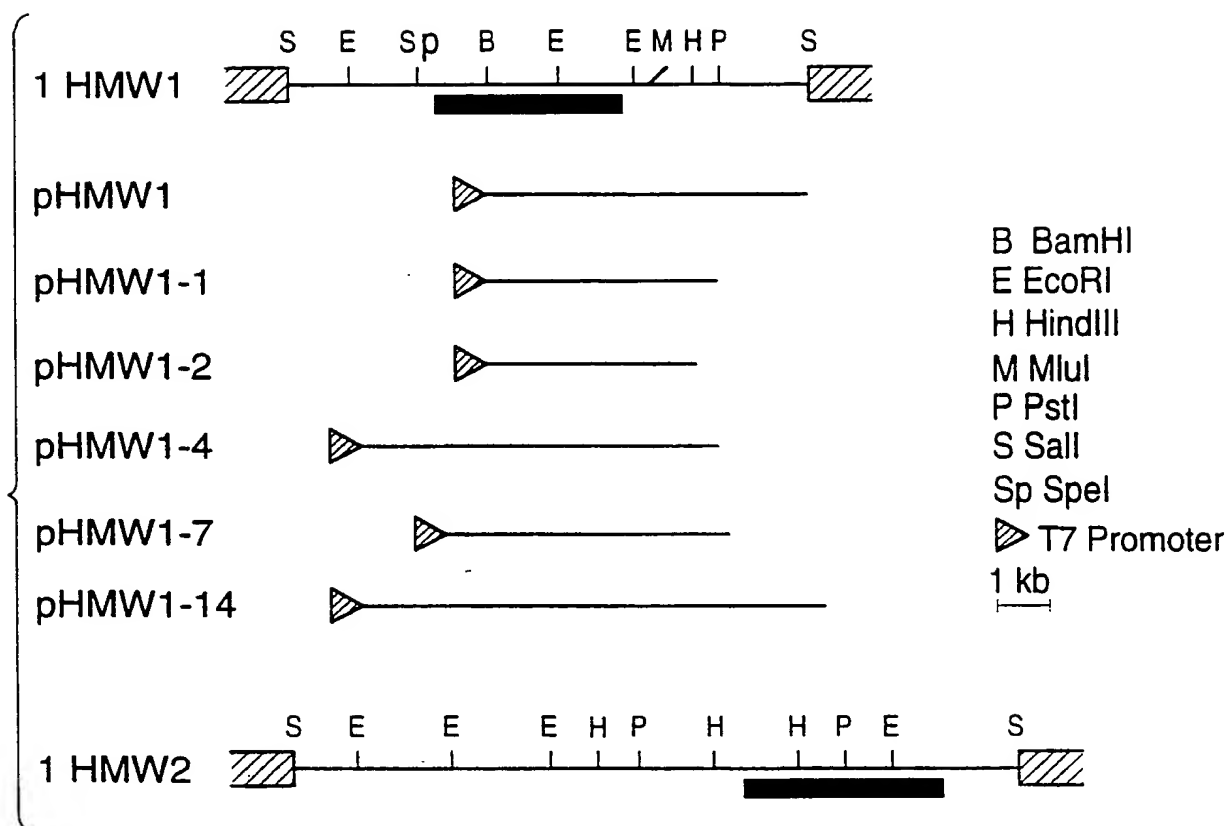
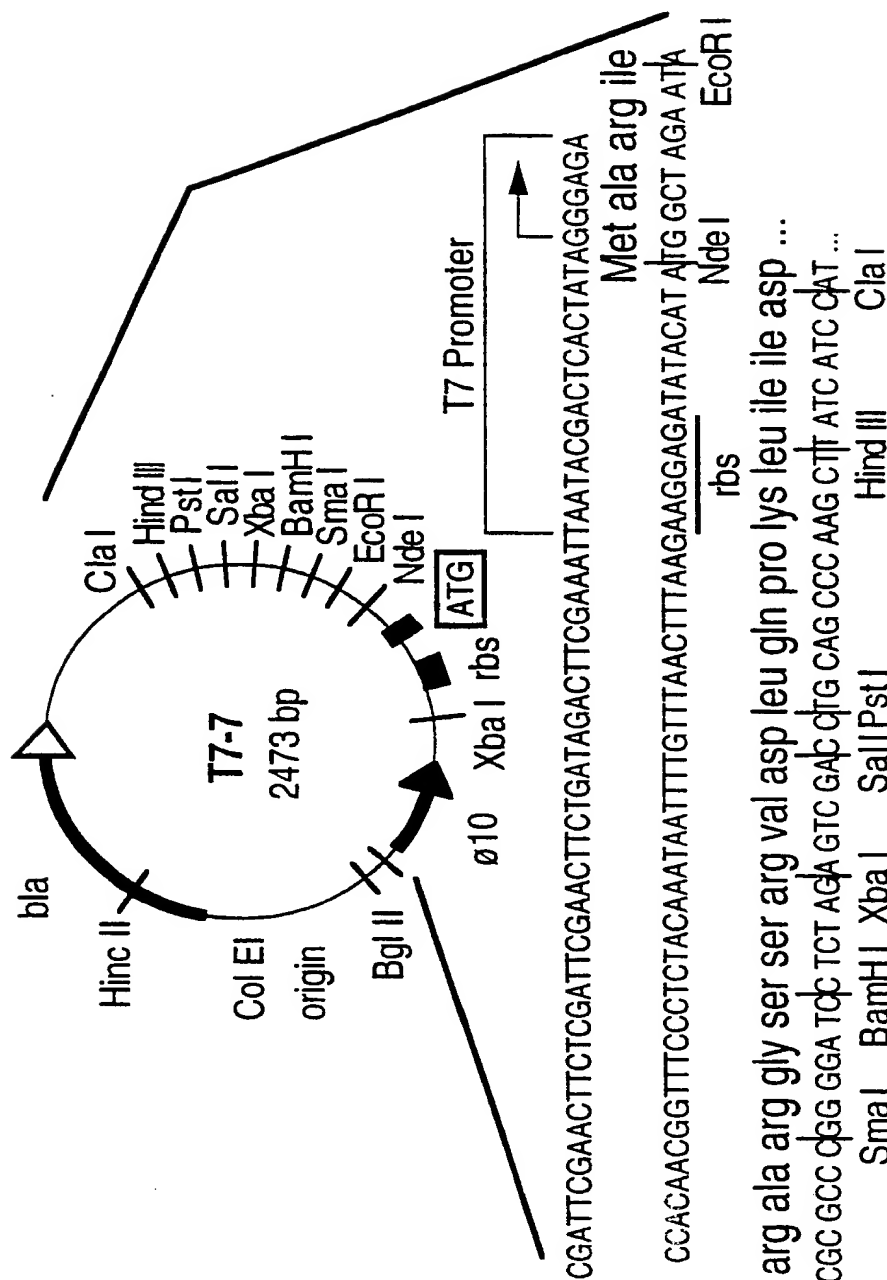


FIG.5 A.

20/68

**FIG. 5B.**

(A) Partial restriction maps of representative HMW1 and HMW2 recombinant phage and of HMW1 plasmid subclones. The shaded boxes indicate the locations of the structural genes. In the recombinant phage, transcription proceeds from left to right for the HMW1 gene and from right to left for the HMW2 gene. The methods used for construction of the plasmids shown are described in the text. (B) Restriction map of the T7 expression vector pT7-7. This vector contains the T7 RNA polymerase promoter $\phi 10$, a ribosome - binding site (rbs), and the translational start site for the T7 gene 10 protein upstream from a multiple cloning site (37).

FIG. 6A.

1 ACAGCGTTCT CTTAATACTA GTACAAACCC ACAATAAAAT ATGACAAACA
 51 ACAATTACAA CACCTTTTTT GCAGTCTATA TGCAAAATATT TTAAAAAATA
 101 GTATAAATCC GCCATATAAA ATGGTATAAT CTTCATCTTT TCATCTTTCA
 151 TCTTTTCATCT TTCATCTTTC ATCTTTTCATC TTTTCATCTTT CATCTTTTCAT
 201 CTTTTCATCTT TCATCTTTCA TCTTTTCATCT TTCATCTTTC ACATGAAATG
 251 ATGAACCGAG GGAAGGGAGG GAGGGCAAG AATGAAGAGG GAGCTGAACG
 301 AACGCAAAATG ATAAAGTAAT TTAATTGTTC AACTAACCTT AGGAGAAAAT
 351 ATGAACAAGA TATATCGTCT CAAATTCAGC AAACGCCCTGA ATGCTTTGGT
 401 TGCTGTGTCT GAATTGGCAC GGGTTGTGA CCATTCCACA GAAAAAGGCA
 451 GCGAAAAACC TGCTCGCATG AAAGTGCCTC ACTTAGCGTT AAAGCCACTT
 501 TCCGCTATGT TACTATCTTT AGGTGTAACA TCTATTCCAC AATCTGTTTT
 551 AGCAAGCGGC TTACAAGGAA TGGATGTAGT ACACGGCACA GCCACTATGC
 601 AAGTAGATGG TAATAAAACC ATTATCCGCA ACAGTGTGTA CGCTATCATT
 651 AATTGGAAAC AATTAAACAT CGACCAAAAT GAAATGGTGC AGTTTTTACA
 701 AGAAAACAAC AACTCCGCCG TATTCAACCG TGTACATCT AACCAATCT
 751 CCCAATTAAA AGGGATTTTA GATCTAACG GACAAGTCTT TTTAATCAAC

21/68

FIG. 6B.

801 CCAAATGGTA TCACAAATAGG TAAAGACGCA ATTATTAACA CTAATGGCTT
851 TACGGGCTTCT ACGCTAGACA TTTCTAACGA AAACATCAAG GCGCGTAATT
901 TCACCTTCGA GCAAACCAA GATAAAGCGC TCGCTGAAAT TGTGAATCAC
951 GGTTTAATTA CTGTCGGTAA AGACGGCAGT GTAAATCTTA TTGGTGGCAA
1001 AGTGAAAAAC GAGGTGTGA TTAGCGTAAA TGGTGGCAGC ATTTCTTTAC
1051 TCGCAGGGCA AAAAATCACC ATCAGCGATA TAATAAACCC AACCATTACT
1101 TACAGCATTG CCGCGCCTGA AAATGAAGCG GTCAATCTGG GCGATATTTT
1151 TGCCAAAGGC GGTAACATTA ATGTCCGTGC TGCCACTATT CGAAACCAAG
1251 CTTTCCGCCA AAGAGGGTGA AGCGGAAATT GGCGGTGTAA TTTCCGCTCA
1301 AAATCAGCAA GCTAAAGGCG GCAAGCTGAT GATTACAGGC GATAAAGTCA
1351 CATTA AAAAC AGGTGCAGTT ATCGACCTTT CAGGTAAGA AGGGGAGAA
1401 ACTTACCTTG GCGGTGACGA GCGGGCGGAA GGTA AAAACG GCATTCAATT
1451 AGCAAAGAAA ACCTCTTTAG AAAAAGGCTC AACCATCAAT GTATCAGGCA
1501 AAGAAAAAGG CGGACGCGCT ATTGTGTGGG GCGATATTGC GTTAATGAC
1551 GGCAATATTA ACGCTCAAGG TAGTGGTGAT ATCGCTAAA CCGGTGGTTT
1601 TGTGAGACG TCGGGGCATG ATTTATTCAT CAAAGACAAT GCAATTGTG

22/08

FIG. 6C.

1651 ACGCCAAAGA GTGGTTGTTA GACCCGGATA ATGTATCTAT TAATGCAGAA
 1701 ACAGCAGGAC GCAGCAATAC TTCAGAAGAC GATGAATACA CGGGATCCGG
 1751 GAATAGTGCC AGCACCCCAA AACGAAACAA AGAAAAGACA ACATTAACAA
 1801 ACACAACTCT TGAGAGTATA CTAAAAAAG GTACCTTTGT TAACATCACT
 1851 GCTAATCAAC GCATCTATGT CAATAGCTCC ATTAATTTAT CCAATGGCAG
 1901 CTTAACTCTT TGGAGTGAGG GTCGGAGCGG TGGCGGCGTT GAGATTAAACA
 1951 ACGATATTAC CACCGGTGAT GATACCAGAG GTGCAAACTT AACAAATTAC
 2001 TCAGGGCGCT GGGTTGATGT TCATAAAAT ATCTCACTCG GGGCGCAAGG
 2051 TAACATAAAC ATTACAGCTA AACAAAGATAT CGCCTTTGAG AAAGGAAGCA
 2101 ACCAAGTCAT TACAGGTCAA GGGACTATTA CCTCAGGCAA TCAAAAAGGT
 2151 TTTAGATTTA ATAATGTCTC TCTAAACGGC ACTGGCAGCG GACTGCAATT
 2201 CACCACTAAA AGAACCAATA AATACGCTAT CACAAATAAA TTTGAAGGGA
 2251 CTTTAAATAT TTCAGGGAAA GTGAACATCT CAATGGTTTT ACCTAAAAAT
 2301 GAAAGTGGAT ATGATAAATT CAAAGGACGC ACTTACTGGA ATTTAACCTC
 2351 GAAAGTGGAT ATGATAAATT CAAAGGACGC CCTCACTATT GACTCCAGAG
 2401 GAAGCGATAG TGCAGGCACA CTTACCCAGC CTTATAAATT AAACGGTATA
 2451 TCATTCAACA AAGACACTAC CTTTAATGTT GAACGAAATG CAAGAGTCAA

23 / 88

FIG. 6D.

2501 CTTTGACATC AAGGCACCAA TAGGGATAAA TAAGTATTCT AGTTTGAATT
2551 ACGCATCATT TAATGGAAAC ATTTCAAGTTT CGGGAGGGGG GAGTGTGAT
2601 TTCACACTTC TCGCCTCATC CTCTAACGTC CAAACCCCCG GTGTAGTTAT
2651 AAATTCTAAA TACTTTAATG TTTCACACAG GTCAAGTTTA AGATTAAAA
2701 CTTCAGGCTC AACAAAACT GGCTTCTCAA TAGAGAAAGA TTTAACTTTA
2751 AATGCCACCG GAGGCAACAT AACACTTTTG CAAGTTGAAG GCACCGATGG
2801 AATGATTGGT AAAGGCATTG TAGCCAAAAA AACATAACC TTTGAAGGAG
2851 GTAAGATGAG GTTTGGCTCC AGGAAAGCCG TAACAGAAAT CGAAGGCAAT
2901 GTTACTATCA ATAACAACGC TAACGTCACT CTTATCGGTT CGGATTTTGA
2951 CAACCATCAA AAACCTTTAA CTATTAAAA AGATGTCATC ATTAATAGCG
3001 GCAACCTTAC CGCTGGAGGC AATATTGTCA ATATAGCCGG AAATCTTACC
3051 GTTGAAAGTA ACGCTAATT CAAAGCTATC ACAAATTCA CTTTAAATGT
3101 AGCGGGCTTG TTTGACAACA AAGGCAATTC AAATATTTC ATTGCCAAAG
3151 GAGGGGCTCG CTTTAAAGAC ATTGATAATT CCAAGAATTT AAGCATCACC
3201 ACCAACTCCA GCTCCACTTA CCGCACTATT ATAAGCGGCA ATATAACCAA
3251 TAAAAACGGT GATTAAATA TTACGAACGA AGGTAGTGAT ACTGAAATGC

24/68

FIG. 6E.

3301	AAATTGGCGG	CGATGTCTCG	CAAAAGAAG	GTAATCTCAC	GATTTCTTCT
3351	GACAAAATCA	ATATTACCA	ACAGATAACA	ATCAAGGCAG	GTGTTGATGG
3401	GGAGAATTCC	GATTCAGACG	CGACAAACAA	TGCCAATCTA	ACCATTAATAA
3451	CCAAGAATT	GAAATTACG	CAAGACCTAA	ATATTTCAGG	TTTCAATAAAA
3501	GCAGAGATTA	CAGCTAAAGA	TGGTAGTGAT	TTAACTATTG	GTAACACCAA
3551	TAGTGCTGAT	GGTACTAATG	CCAAAAAAGT	AACCTTTAAC	CAGGTTAAAG
3601	ATTCAAAAAT	CTCTGCTGAC	GGTCACAAGG	TGACACTACA	CAGCAAAAGTG
3651	GAAACATCCG	GTAGTAATAA	CAACACTGAA	GATAGCAGTG	ACAATAATGC
3701	CGGCTTAACT	ATCGATGCAA	AAAATGTAAAC	AGTAAACAAC	AAATTTACTT
3751	CTCACAAAGC	AGTGAGCATC	TCTGCGACAA	GTGGAGAAAT	TACCACTAAA
3801	ACAGGTACAA	CCATTAAACG	AACCACTGGT	AACGTGGAGA	TAACCGCTCA
3851	AACAGGTAGT	ATCCTAGGTG	GAATTGAGTC	CAGCTCTGGC	TCTGTAAACAC
3901	TTACTGCAAC	CGAGGGCGCT	CTTGCTGTAA	GCAATATTTC	GGGCAACACC
3951	GTTACTGTTA	CTGCAAAATAG	CGGTGCATTA	ACCACTTTGG	CAGGCTCTAC
4001	AATTAAAGGA	ACCGAGAGTG	TAACCACTTC	AAGTCAATCA	GGCGATATCG
4051	GCGGTACGAT	TTCTGGTGGC	ACAGTAGAGG	TTAAAGCAAC	CGAAAGTTTA

FIG. 6F.

4101 ACCACTCAAT CCAATTCAAA AATTAAAGCA ACAACAGGCG AGGCTAACGT
4151 AACAAGTGCA ACAGGTACAA TTGGTGGTAC GATTTCGGT AATACGGTAA
4201 ATGTTACGGC AAACGCTGGC GATTTAACAG TTGGGAATGG CGCAGAAATT
4251 AATGCGACAG AAGGAGCTGC AACCTTAACT ACATCATCGG GCAAATTAAC
4301 TACCGAAGCT AGTTCACACA TTACTTCAGC CAAGGGTCAG GTAAATCTTT
4351 CAGCTCAGGA TGGTAGCGTT GCAGGAAGTA TTAATGCCGC CAATGTGACA
4401 CTAAATACTA CAGGCACTTT AACTACCGTG AAGGGTTCAA ACATTAATGC
4451 AACCAGCGGT ACCTTGGTTA TTAACGCAA AGACGCTGAG CTAAATGGCG
4501 CAGCATTTGG TAACCAACACA GTGGTAAATG CAACCAACGC AAATGGCTCC
4551 GGCAGCGTAA TCGCGACAAC CTCAAGCAGA GTGAACATCA CTGGGGATTT
4601 AATCACAAATA AATGGATTAA ATATCATTTT AAAAAACGGT ATAAACACCG
4651 TACTGTTAAA AGGCGTTAAA ATTGATGTGA AATACATTCA ACCGGGTATA
4701 GCAAGCGTAG ATGAAGTAAT TGAAGCGAAA CGCATCCTTG AGAAGGTAAA
4751 AGATTTATCT GATGAAGAAA GAGAAGCGTT AGCTAAACTT GGCCTAAGTG
4801 CTGTACGTTT TATTGAGCCA AATAATACAA TTACAGTCGA TACACAAAAT
4851 GAATTTGCAA CCAGACCATT AAGTCGAATA GTGATTTCTG AAGCAGGGC
4901 GTGTTTCTCA AACAGTGATG GCGCGACGGT GTGCGTTAAT ATCGCTGATA

FIG. 6G.

4951	ACGGGCGGTA	GCGTCAGTA	ATTGACAAGG	TAGATTTTCAT	CCTGCAATGA
5001	AGTCATTTTA	TTTTCGTATT	ATTTACTGTG	TGGGTTAAAG	TTCAGTACGG
5051	GCTTTACCCA	TCTTGTA AAA	AATTACGGAG	AATACAATAA	AGTATTTTTA
5101	ACAGGTTATT	ATTATGAAAA	ATATAAAAAG	CAGATTA AAA	CTCAGTGCAA
5151	TATCAGTATT	GCTTGGCCTG	GCTTCTTCAT	CATTGTATGC	AGAAGAAGCG
5201	TTTTTTAGTAA	AAGGCTTTCA	GTTATCTGGT	GCAC TTGAAA	CTTTAAGTGA
5251	AGACGCCCAA	CTGTCGTAG	CAAAATCTTT	ATCTAAATAC	CAAGGCTCGC
5301	AAACTTTAAC	AAACCCTAAA	ACAGCACAGC	TTGAATTACA	GGCTGTGCTA
5351	GATAAGATTG	AGCCAAAATA	GTTTGATGTG	ATATTGCCAC	AACAAAACCAT
5401	TACGGATGGC	AATATTATGT	TTGAGCTAGT	CTCGAAATCA	GCCGCAGAAA
5451	GCCAAGTTTT	TTATAAGGCG	AGCCAGGGTT	ATAGTGAAGA	AAATATCGCT
5501	CGTAGCCCTGC	CATCTTTGAA	ACAAGGAAAA	GTGTATGAAG	ATGGTCGTCA
5551	GTGGTTCGAT	TTGCGTGAAT	TCAATATGGC	AAAAGAAAAT	CCACTTAAAG
5601	TCACTCGCGT	GCATTACGAG	TTAAACCCCTA	AAAACAAAAC	CTCTGATTTG
5651	GTAGTTGCAG	GTTTTTCGCC	TTTTTGGCAA	ACGCGTAGCT	TTGT TTCCTA
5701	TGATAAATTTC	GGCGCAAGGG	AGTTTAACTA	TCAACGTGTA	AGCTAGGTT

27/88

FIG. 6H.

5751 TTGTAATGC CAATTGACC GGACATGATG ATGTATTAAA TCTAAACGCA
5801 TTGACCAATG TAAAGCACC ATCAAAATCT TATGCGGTAG GCATAGGATA
5851 TACTTATCCG TTTTATGATA AACACCAATC CTTAAGTCTT TATACCAGCA
5901 TGAGTTATGC TGATTCTAAT GATATCGACG GCTTACCAAG TCGGATTAAAT
5951 CGTAAATTAT CAAAAGGTCA ATCTATCTCT GCGAATCTGA AATGGAGTTA
6001 TTATCTCCCG ACATTTAACC TTGGAATGGA AGACCAGTTT AAAATTAATT
6051 TAGGCTACAA CTACCGCCAT ATTAATCAAA CATCCGAGTT AACACCCCTG
6101 GGTGCAACGA AGAAAAAATT TGCAGTATCA GCGTAAGTG CAGGCATTGA
6151 TGGACATATC CAATTTACCC CTAAAACAAT CTTAATATT GATTAACTC
6201 ATCATTATTA CGCGAGTAAA TTACCAGGCT CTTTGTGAAT GGAGCGCATT
6251 GGCGAAACAT TTAATCGCAG CTATCACATT AGCACAGCCA GTTTAGGGTT
6301 GAGTCAAGAG TTTGCTCAAG GTTGGCATTT TAGCAGTCAA TTATCGGGTC
6351 AGTTTACTCT ACAAGATATA AGTAGCATAG ATTTATTCTC TGTAACAGGT
6401 ACTTATGGCG TCAGAGGCTT TAAATACGGC GGTGCAAGTG GTGAGCGCGG
6451 TCTTGATATGG CGTAATGAAT TAAAGTATGCC AAAATACACC CGCTTTCAAA
6501 TCAGCCCTTA TGCGTTTTAT GATGCAGGTC AGTTCCGTTA TAATAGCGAA
6551 AATGCTAAAA CTTACGGCGA AGATATGCAC ACGGTATCCT CTGCGGGTTT

20/60

FIG. 6I.

6601	AGGCATTAAA	ACCTCTCCTA	CACAAAACCTT	AAGCTTAGAT	GCTTTTGTG
6651	CTCGTCGCTT	TGCAAAATGCC	AATAGTGACA	ATTGAATGG	CAACAAAAAA
6701	CGCACAAAGCT	CACCTACAAC	CTTCTGGGT	AGATTAAACAT	TCAGTTTCTA
6751	ACCCGTGAAAT	TTAATCAACT	GGTAAGCGTT	CCGCCTACCA	GTTTATAACT
6801	ATATGCTTTA	CCCGCCAATT	TACAGTCTAT	ACGCAACCCT	GTTTTCATCC
6851	TTATATATCA	AACAAACTAA	GCAAACCAAG	CAAACCAAGC	AAACCAAGCA
6901	AACCAAGCAA	ACCAAGCAAA	CCAAGCAAAC	CAAGCAAACC	AAGCAAACCA
6951	AGCAAACCAA	GCAAACCAAG	CAAACCAAGC	AAACCAAGCA	ATGCTAAAAA
7001	ACAAATTTATA	TGATAAACTA	AAACATACTC	CATACCATGG	CAATACAAGG
7051	GATTTAATAA	TATGACAAAA	GAAAATTTAC	AAAGTGTTC	ACAAAATACG
7101	ACCGCTTCAC	TTGTAGAATC	AAACAACGAC	CAAACCTCCC	TGCAAAATACT
7151	TAAACAACCA	CCCAAACCCA	ACCTATTACG	CCTGGAACAA	CATGTCGCCA
7201	AAAAAGATTA	TGAGCTTGCT	TGCCGCGAAT	TAAATGGCGAT	TTTGGAAAAA
7251	ATGGACGCTA	ATTTTGGAGG	CGTTCACGAT	ATTGAATTG	ACGCACCTGC
7301	TCAGCTGGCA	TATCTACCCG	AAAACTACT	AATTCATTTT	GCCACTCGTC
7351	TCGCTAATGC	AATTACAACA	CTCTTTTCCG	ACCCCGAATT	GGCAATTTC

FIG. 6J.

7401	GAAGAAGGG	CATTAAAGAT	GATTAGCCTG	CAACGCTGGT	TGACGCTGAT
7451	TTTTGCCCTCT	TCCCCCTACG	TTAACGCAGA	CCATATTCTC	AATAAATATA
7501	ATATCAACCC	AGATTCCGAA	GGTGGCTTTC	ATTTAGCAAC	AGACAACTCT
7551	TCTATTGCTA	AATTCTGTAT	TTTTTACTTA	CCCGAATCCA	ATGTCAATAT
7601	GAGTTTAGAT	GCGTTATGGG	CAGGGAATCA	ACAACTTTGT	GCTTCATTGT
7651	GTTTTGCGTT	GCAGTCTTCA	CGTTTTATTG	GTA CTGCATC	TGCGTTTCAT
7701	AAAAGAGCGG	TGGTTTTACA	GTGGTTTCCT	AAAAAACTCG	CCGAAAATTGC
7751	TAATTTAGAT	GAATTGCCTG	CAAATATCCT	TCA TGATGTA	TATATGCACT
7801	GCAGTTATGA	TTTAGCAAAA	AACAAGCACG	ATGTTAAGCG	TCCATTAAAC
7851	GAACTTGTC	GCAAGCATAT	CCTCACGCAA	GGATGGCAAG	ACCGCTACCT
7901	TTACACCTTA	GGTAAAAAGG	ACGGCAAACC	TGTGATGATG	GTA CTGCTTG
7951	AACATTTTAA	TTCGGGACAT	TCGATTTATC	GCACGCATTC	AAC TTCAATG
8001	ATTGCTGCTC	GAGAAAAATT	CTATTTAGTC	GGCTTAGGCC	ATGAGGGCGT
8051	TGATAACATA	GGTCGAGAAG	TGTTTGACGA	GTTCTTTGAA	ATCAGTAGCA
8101	ATAATATAAT	GGAGAGACTG	TTTTTTTATCC	GTA AACAGTG	CGAAACTTTC
8151	CAACCCGCAG	TGTTCTATAT	GCCAAGCATT	GGCATGGATA	TTACCACGAT

FIG. 6K.

8201 TTTTGTGAGC AACACTCGGC TTGCCCCCTAT TCAAGCTGTA GCCTTGGGTC
 8251 ATCCTGCCAC TACGCATTCT GAATTTATTG ATTATGTCAT CGTAGAAGAT
 8301 GATTATGTGG GCAGTGAAGA TTGTTTAGC GAAACCCCTTT TACGCTTACC
 8351 CAAAGATGCC CTACCTTATG TACCATCTGC ACTCGCCCCCA CAAAAAGTGG
 8401 ATTATGTACT CAGGGAAAAC CCTGAAGTAG TCAATATCGG TATTGCCGCT
 8451 ACCACAAATGA AATTAAACCC TGAATTTTGG CTAACATTCG AAGAAATCAG
 8501 AGATAAAGCT AAAGTCAAAA TACATTTTCA TTTCGCACCT GGACAAATCAA
 8551 CAGGCTTGAC ACACCCCTTAT GTCAAATGGT TTATCGAAG CTATTTAGGT
 8601 GACGATGCCA CTGCACATCC CCACGCACCT TATCACGATT ATCTGGCAAT
 8651 ATTGCGTGAT TCGGATATGC TACTAAATCC GTTTCCTTTC GGTAATACTA
 8701 ACGGCATAAT TGATATGGTT ACATTAGGTT TAGTTGGTGT ATGCAAAACG
 8751 GGGATGAAG TACATGAACA TATTGATGAA GGTCTGTTTA AACGCTTAGG
 8801 ACTACCAGAA TGGCTGATAG CCGACACACG AGAAACATAT ATTGAATGTG
 8851 CTTTGCGTCT AGCAGAAAAC CATCAAGAAC GCCTTGAACT CCGTCGTTAC
 8901 ATCATAGAAA ACAACGGCTT ACAAAAAGCTT TTTACAGGCG ACCCTCGTCC
 8951 ATTGGGCAA ATACTGCTTA AGAAAACAAA TGAATGGAAG CGGAAGCACT
 9001 TGAGTAAAAA ATAACGGTTT TTTAAAGTAA AAGTCCGGTT AATTTTCAA

31/68

32 / 68

FIG. 6L.

9051	GCGTTTTAAA	AACCTCTCAA	AAATCAACCG	CACTTTTATC	TTTATAACGC
9101	TCCCGCGCGC	TGACAGTTTA	TCTCTTCTT	AAAATACCCA	TAAAATTGTG
9151	GCAATAGTTG	GGTAATCAAA	TTCAATTGTT	GATACGGCAA	ACTAAAGACG
9201	GCGCGTTCTT	CGGCAGTCAT	C		

FIG. 7A.

1 CGCCACTTCA ATTTTGGATT GTTGAAATTC AACTAACCAA AAAGTGCGGT
 51 TAAAAATCTGT GGAGAAAATA GGTGTAGTG AAGAACGAGG TAAATTGTTCA
 101 AAAGGATAAA GCTCTCTTAA TTGGGCATTG GTTGGCGTTT CTTTTCGGT
 151 TAAATAGTAA TTATATTCTG GACGACTATG CAATCCACCA ACAACTTTAC
 201 CGTTGGTTTT AAGCGTTAAT GTAAGTTCTT GCTCTTCTTG GCGAATACGT
 251 AATCCCATTT TTTGTTTAGC AAGAAAATGA TCGGGATAAT CATAATAGGT
 301 GTTGCCCCAA AATAAATTTT GATGTTCTAA AATCATAAAT TTTGCAAGAT
 33 / 68
 351 ATTGTGGCAA TTCAATACCT ATTTGTGGCG AAATCGCCAA TTTTAATTCA
 401 ATTTCTTGTA GCATAATATT TCCCACCTCA ATCAACTGGT TAAATATACA
 451 AGATAATAAA AATAAATCAA GATTTTGTG ATGACAAACA ACAATTACAA
 501 CACCTTTTTT GCAGTCTATA TGCAAAATAT TTAAAAAAAT AGTATAAATC
 551 CGCCATATAA AATGGTATAA TCTTTCATCT TTTCATCTTTC ATCTTTCATC
 601 TTTTCATCTTT CATCTTTTCAT CTTTTCATCTT TCATCTTTCA TCTTTCATCT
 651 TTTCATCTTTC ATCTTTTCATC TTTTCATCTTT CACATGAAAT GATGAACCGA
 701 GGAAGGGAG GGAGGGGCAA GAATGAAGAG GGAGCTGAAC GAACGCAAT
 751 GATAAAGTAA TTTAATTGTT CAACTAACCT TAGGAGAAA TATGAACAAG

FIG. 7B.

801	ATATATCGTC	TCAAATTCAG	CAAACGCCCTG	AATGCTTTGG	TTGCTGTGTC
851	TGAATTGGCA	CGGGTTGTG	ACCATTCAC	AGAAAAAGGC	AGCGAAAAAC
901	CTGCTCGCAT	GAAAGTGCCT	CACTTAGCGT	TAAAGCCACT	TTCCGCTATG
951	TTACTATCTT	TAGGTGTAAC	ATCTATTCCA	CAATCTGTTT	TAGCAAGCGG
1001	CAATTTAACA	TCGACCAAAA	TGAAATGGTG	CAGTTTTCAC	AAGAAAAACA
1051	GTAATAAAAC	CATTATCCGC	AACAGTGTG	ACGCTATCAT	TAATTGGAAA
1101	CAATTTAACA	TCGACCAAAA	TGAAATGGTG	CAGTTTTCAC	AAGAAAAACA
1151	CAACTCCGCC	GTATTCAACC	GTGTACATC	TAACCAATC	TCCCAATTAA
1201	AAGGGATTTT	AGATTCTAAC	GGACAAGTCT	TTTTAATCAA	CCCAAAATGGT
1251	ATCACAAATAG	GTAAGACGC	AATTATTAAC	ACTAATGGCT	TTACGGCTTC
1301	TACGCTAGAC	ATTTCTAACG	AAAACATCAA	GGCGCGTAAT	TTCACCCTTCCG
1351	AGCAAACCAA	AGATAAAGCG	CTCGCTGAAA	TTGTGAATCA	CGGTTTAATT
1401	ACTGTCGGTA	AAGACGGCAG	TGTAAATCTT	ATTGGTGGCA	AAGTGAAAAA
1451	CGAGGGGTGTG	ATTAGCGTAA	ATGGTGGCAG	CATTCTTTTA	CTCGCAGGGC
1501	AAAAAATCAC	CATCAGCGAT	ATAATAAACC	CAACCATTAC	TTACAGCATT
1551	GCCGCGCCTG	AAAATGAAGC	GGTCAATCTG	GGCGATATTT	TTGCCAAAGG

FIG. 7C.

1601 CCGTAACATT AATGTCCGTG CTGCCACTAT TCGAAACCAA GTAAACTTT
 1651 CTGCTGATTC TGTAAGCAAA GATAAAGCG GCAATATTGT TCTTTCCGCC
 1701 AAAGAGGGTG AAGCGGAAAT TGGCGGTGTA ATTTCCGCTC AAAATCAGCA
 1751 AGCTAAAGGC GGCAAGCTGA TGATTACAGG CGATAAAGTC ACATTAAAAA
 1801 CAGGTGCAGT TATCGACCCTT TCAGGTAAGG AAGGGGAGA AACTTACCCTT
 1851 GCGGGTGACG AGCGGGCGA AGGTAAAAC GGCATTCAAT TAGCAAAAGAA
 1901 AACCTCTTTA GAAAAAGGCT CAACCATCAA TGTATCAGGC AAAGAAAAAG
 1951 GCGGACGCGC TATTGTGTGG GCGATATTG CGTTAATTGA CGGCAATATT
 2001 AACGCTCAAG GTAGTGGTGA TATCGCTAAA ACCGGTGGTT TTGTGGAGAC
 2051 ATCGGGGCAT TATTTATCCA TTGACAGCAA TGCAATTGTT AAAACAAAAG
 2101 AGTGGTTGCT AGACCCCTGAT GATGTAACAA TTGAAGCCGA AGACCCCTT
 2151 CGCAATAATA CCGGTATAAA TGATGAATTC CCAACAGGCA CCGGTGAAGC
 2201 AAGCGACCCT AAAAAAATA GCGAACTCAA AACACGCTA ACCAATACAA
 2251 CTATTTCAA TTAATCTGAAA AACGCCCTGGA CAATGAATAT AACGGCATCA
 2301 AGAAAACTTA CCGTTAATAG CTCAATCAAC ATCGGAAGCA ACTCCCCTT
 2351 AATTCTCCAT AGTAAAGGTC AGCGTGCGG AGGCGTTCAG ATTGATGGAG
 2401 ATATTACTTC TAAAGCGGA AATTTAACCA TTTATTCTGG CGGATGGGTT

FIG. 7D.

2451	GATGTTTCATA	AAAATATTAC	GCTTGATCAG	GGTTTTTTTAA	ATATTACCGC
2501	CGCTTCCCGTA	GCTTTTGAAG	GTGGAATAA	CAAAGCACGC	GACGCGGCAA
2551	ATGCTAAAT	TGTCGCCCAG	GGCACTGTAA	CCATTACAGG	AGAGGGAAAA
2601	GATTTCAGGG	CTAACAACGT	ATCTTTAAAC	GGAACGGGTA	AAGGTCTGAA
2651	TATCATTTCA	TCAGTGAATA	ATTTAACCCA	CAATCTTAGT	GGCACAATTA
2701	ACATATCTGG	GAATATAACA	ATTAACCAA	CTACGAGAAA	GAACACCTCG
2751	TATTGGCAA	CCAGCCATGA	TTCGCACTGG	AACGTCAGTG	CTCTTAATCT
2801	AGAGACAGGC	GCAAATTTTA	CCTTTATTAA	ATACATTTCA	AGCAATAGCA
2851	AAGGCTTAAC	AACACAGTAT	AGAAGCTCTG	CAGGGGTGAA	TTTTAACGGC
2901	GTAAATGGCA	ACATGTCAAT	CAATCTCAA	GAAGGAGCGA	AAGTTAATTT
2951	CAAATTAAAA	CCAAACGAGA	ACATGAACAC	AAGCAAACCT	TTACCAATTC
3001	GGTTTTTAGC	CAATATCACA	GCCACTGGTG	GGGGCTCTGT	TTTTTTTGAT
3051	ATATATGCCA	ACCATTCTGG	CAGAGGGGCT	GAGTTAAAAA	TGAGTGAAAT
3101	TAATATCTCT	AACGGCGCTA	ATTTTACCCT	AAATTCCCAT	GTTTCGGCGG
3151	ATGACGCTTT	TAAATCAAC	AAAGACTTAA	CCATAAATGC	AACCAATTCA
3201	AATTTCAGCC	TCAGACAGAC	GAAAGATGAT	TTTTTATGACG	GGTACGCACG

FIG. 7E.

3251 CAATGCCATC AATTCAACCT ACAACATATC CATTCTGGGC GGTAATGTCA
 3301 CCCTTGGTGG ACAAAACTCA AGCAGCAGCA TTACGGGGAA TATTACTATC
 3351 GAGAAAGCAG CAAATGTTAC GCTAGAAGCC AATAACGCC CTAATCAGCA
 3401 AAACATAAGG GATAGAGTTA TAAAACTTGG CAGCTTGCTC GTTAATGGGA
 3451 GTTTAAGTTT AACTGGCGAA AATGCAGATA TTAAAGGCAA TCTCACTATT
 3501 TCAGAAAGCG CCACTTTTAA AGGAAAGACT AGAGATACCC TAAATATCAC
 3551 CGGCAATTTT ACCAATAATG GCACTGCCGA AATTAATATA ACACAAGGAG
 3601 TGGTAAAAC TGGCAATGTT ACCAATGATG GTGATTTAAA CATTACCACT
 3651 CACGCTAAAC GCAACCAAAG AAGCATCATC GCGGGAGATA TAATCAACAA
 3701 AAAAGGAAGC TTAAATATTA CAGACAGTAA TAATGATGCT GAAATCCAAA
 3751 TTGGCGGCAA TATCTCGCAA AAAGAAGGCA ACCTCACGAT TTCCTTCCGAT
 3801 AAAATTAAATA TCACCAAACA GATAACAATC AAAAAGGGTA TTGATGGAGA
 3851 GGACTCTAGT TCAGATGCGA CAAGTAATGC CAACCTAACT ATTAAAACCA
 3901 AAGAATTGAA ATTGACAGAA GACCTAAGTA TTTTCAGGTTT CAATAAAGCA
 3951 GAGATTACAG CCAAAGATGG TAGAGATTTA ACTATTGGCA ACAGTAATGA
 4001 CGGTAACAGC GGTGCCGAAG CCAAACAGT AACTTTTAAC AATGTTAAAG

37/68

FIG. 7F.

4051 ATTCAAAAAT CTCTGCTGAC GGTCACAATG TGACACTAAA TAGCAAAGTG
4101 AAAACATCTA GCAGCAATGG CGGACGTGAA AGCAATAGCG ACAACGATAC
4151 CGGCTTAACT ATTACTGCAA AAAATGTAGA AGTAAACAAA GATATTACTT
4201 CTCTCAAAAC AGTAAATATC ACCGCGTCGG AAAAGGTTAC CACCACAGCA
4251 GGCTCGACCA TTAACGCAAC AAATGGCAAA GCAAGTATTA CAACCAAAAC
4301 AGGTGATATC AGCGGTACGA TTTCCGGTAA CACGGTAAGT GTTAGCGCGA
4351 CTGGTGATTT AACCACTAAA TCCGGCTCAA AAATTGAAGC GAAATCGGGT
4401 GAGGCTAATG TAACAAGTGC AACAGGTACA ATTGGCGGTA CAATTTCCGG
4451 TAATACGGTA AATGTTACGG CAAACGCTGG CGATTTAACA GTTGGGAATG
4501 GCGCAGAAAT TAATGCGACA GAAGGAGCTG CAACCTTAAC CGCAACAGGG
4551 AATACCTTGA CTACTGAAGC CGGTTCTAGC ATCACTTCAA CTAAGGGTCA
4601 GGTAGACCTC TTGGCTCAGA ATGGTAGCAT CGCAGGAAGC ATTAATGCTG
4651 CTAATGTGAC ATTAAATACT ACAGGCACCT TAACCAACCGT GGCAGGCTCG
4701 GATATTAAAG CAACCAGCGG CACCTTGGTT ATTAACGCAA AAGATGCTAA
4751 GCTAAATGGT GATGCATCAG GTGATAGTAC AGAAGTGAAT GCAGTCAACG
4801 ACTGGGGATT TGGTAGTGTG ACTGCGGCAA CCTCAAGCAG TGTGAATATC
4851 ACTGGGGATT TAAACACAGT AAATGGGTTA AATATCATTT CGAAAGATGG

38/68

FIG. 7G.

4901 TAGAAACACT GTGCGCTTAA GAGCAAGGA AATTGAGGTG AAATATATCC
 4951 AGCCAGGTGT AGCAAGTGTA GAAGAAGTAA TTGAAGCGAA ACGCGTCCTT
 5001 GAAAAAGTAA AAGATTATC TGATGAAGAA AGAGAAACAT TAGCTAAACT
 5051 TGGTGTAAGT GCTGTACGTT TTGTTGAGCC AAATAATACA ATTACAGTCA
 5101 ATACACAAAA TGAATTTACA ACCAGACCGT CAAGTCAAGT GATAATTTCT
 5151 GAAGGTAAGG CGTGTTTCTC AAGTGTAAT GCGGCACGAG TATGTACCAA
 5201 TGTGTGCTGAC GATGGACAGC CGTAGTCAGT AATTGACAAAG GTAGATTTC³
 5251 TCCTGCAATG AAGTCATTTT ATTTTCGTAT TATTTACTGT GTGGGTTAAA³
 5301 GTTCAGTACG GGCTTTACCC ATCTTGTA³
 5351 AAGTATTTTT AACAGGTAT TATTATGAAA AATATAAAAA GCAGATTAAA³
 5401 ACTCAGTGCA ATATCAGTAT TGCTTGGCCT GGCTTCTTCA TCATTGTATG
 5451 CAGAAGAAGC GTTTT³TAGTA AAAGGCTTTC AGTTATCTGG TGCACTTGAA
 5501 ACTTTAAGTG AAGACGCCCA ACTGTCTGTA GCAAAAATCTT TATCTAAATA
 5551 CCAAGGCTCG CAAACTTTAA CAAACCTAAA AACAGCACAG CTTGAATTAC
 5601 AGGCTGTGCT AGATAAGATT GAGCCAAAATA AATTGATGT GATATTGCCG
 5651 CAACAAACCA TTACGGATGG CAATATCATG TTTGAGCTAG TCTCGAAATC

FIG. 7H.

5701	AGCCGCAGAA	AGCCAAGTTT	TTTATAAGGC	GAGCCAGGGT	TATAGTGAAG
5751	AAAATATCGC	TCGTAGCCCTG	CCATCTTTGA	AACAAGGAAA	AGTGTATGAA
5801	GATGGTCGTC	AGTGGTTCGA	TTTGCGTGAA	TTTAATATGG	CAAAAGAAAA
5851	CCCGCTTAAG	GTTACCCCGTG	TACATTACGA	ACTAAACCCCT	AAAAACAAAA
5901	CCTCTAATT	GATAATTGCG	GGCTTCTCGC	CTTTTGGTAA	AACGCGTAGC
5951	TTTATTTCCT	ATGATAATT	CGGCGCGAGA	GAGTTTAACT	ACCAACGTGT
6001	AAGCTTGGGT	TTTGTTAATG	CCAATTTAAC	TGGTCATGAT	GATGTGTTAA
6151	TTATACCAGT	ATGAGTTATG	CTGATTCTAA	TGATATCGAC	GGCTTACCAA
6201	GTGCGATTAA	TCGTAATAA	TCAAAAGGTC	AATCTATCTC	TGCGAATCTG
6251	AAATGGAGTT	ATTATCTCCC	AACATTTAAC	CTTGGCATGG	AAGACCAATT
6301	TAAAATTAAT	TTAGGCTACA	ACTACCGCCA	TATTAAATCAA	ACCTCCGCGT
6351	TAAATCGCTT	GGGTGAAACG	AAGAAAAAAT	TTGCAGTATC	AGGCGTAAAGT
6401	GCAGGCATTG	ATGGACATAT	CCAATTTACC	CCTAAAACAA	TCTTTAATAT
6451	TGATTTAACT	CATCATTTAT	ACGCGAGTAA	ATTACCAGGC	TCTTTTGGAA
6501	TGGAGCGCAT	TGGCGAAACA	TTTAATCGCA	GCTATCACAT	TAGCACAGCC
6551	AGTTTAGGGT	TGAGTCAAGA	GTTTGCTCAA	GGTTGGCAT	TTAGCAGTCA
6601	ATTATCAGGT	CAATTTACTC	TACAAGATAT	TAGCAGTATA	GATTATTCT

40/68

FIG. 7I.

6651 CTGTAACAGG TACTTATGGC GTCAGAGGCT TTAAATACGG CCGTGCAAGT
 6701 GGTGAGCGCG GTCTTGTATG GCGTAATGAA TTAAAGTATGC CAAAATACAC
 6751 CCGCTTCCAA ATCAGCCCTT ATGCGTTTAA TGATGCAGGT CAGTCCGTT
 6801 ATAATAGCGA AAATGCTAAA ACTTACGGCG AAGATATGCA CACGGTATCC
 6851 TCTGCGGGTT TAGGCATTAA AACCTCTCCT ACACAAAAC TAAAGCCTAGA
 6901 TGCTTTTGTG GCTCGTCGCT TTGCAAAATG CAATAGTGAC AATTGAATG
 6951 GCAACAAAAA ACGCACAGC TCACCTACAA CCTTCTGGGG GAGATTAACA
 7001 TTCAGTTTCT AACCCGTGAA TTTAATCAAC TGGTAAGCGT TCCGCCCTACC
 7051 AGTTTATAAC TATATGCTTT ACCCGCCAAT TTACAGTCTA TAGGCAACCC
 7101 TGTTTTTACC CTTATATATC AAATAAACAA GCTAAGCTGA GCTAAGCAAA
 7151 CCAAGCAAAC TCAAGCAAGC CAAGTAATAC TAAAAAACA ATTTATATGA
 7201 TAAACTAAAG TATACTCCAT GCCATGGCGA TACAAGGGAT TTAATAATAT
 7251 GACAAAAGAA AATTTGCAA ACGCTCCTCA AGATGCGACC GCTTACTTG
 7301 CGGAATTAA GCAACAATCA ACTCCCCTGC GAATATTAA ACAACCACGC
 7351 AAGCCCAGCC TATTACGCTT GGAACAACAT ATCGCAAAA AAGATTATGA
 7401 GTTTGCTTGT CGTGAATTAA TGGTGATTCT GGAATAAATG GACGCTAATT

41/88

FIG. 7J.

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7451 TTGGAGGCGT TCACGATATT GAATTGACG CACCCGCTCA GCTGGCATAT
7501 CTACCCGAAA AATTACTAAT TTATTTTGCC ACTCGTCTCG CTAATGCAAT
7551 TACAACACTC TTTTCCGACC CCGAATTGGC AATTCTGAA GAAGGGCGGT
7601 TAAAGATGAT TAGCCTGCAA CGCTGGTTGA CGTGATTTT TGCCTCTTCC
7651 CCTACGTTA ACGCAGACCA TATTCTCAAT AAATAATAA TCAACCCAGA
7701 TTCCGAAGGT GGCTTTCATT TAGCAACAGA CAACTCTTCT ATTGCTAAAT
7751 TCTGTATTTT TTACTIONACC GAATCCAATG TCAATATGAG TTTAGATGCG
7801 TTATGGGCAG GGAATCAACA ACTTTGTGCT TCATTGTGTT TTGCGTTGCA
7851 GTCCTCACGT TTTATTGGTA CCGCATCTGC GTTTCATAAA AGAGCGGTGG
7901 TTTTACAGTG GTTCCCTAAA AACTCGCCG AAATTGCTAA TTTAGATGAA
7951 TTGCCCTGCAA ATATCCTTCA TGATGTATAT ATGCACCTGCA GTTATGATTT
8001 AGCAAAAAC AAGCACGATG TTAAGCGTCC ATTAAACGAA CTGTGTCGCA
8051 AGCATATCCT CACGCAAGGA TGGCAAGACC GCTACCCTTA CACCTTAGGT
8101 AAAAAGGACG GCAAACCTGT GATGATGGTA CTGCTTGAAC ATTTTAATTC
8151 GGGACATTTC ATTTATCGTA CACATCAAC TTCAATGATT GCTGCTCGAG
8201 AAAAATTCTA TTTAGTCGGC TTAGGCCCATG AGGCGGTTGA TAAAATAGGT

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FIG. 7K.

8251 CGAGAAGTGT TTGACGAGTT CTTTGAAATC AGTAGCAATA ATATAATGGA
 8301 GAGACTGTTT TTTATCCGTA AACAGTGCGA AACTTTCCAA CCCGCAGTGT
 8351 TCTATATGCC AAGCATTTGGC ATGGATATTA CCACGATTTT TGTGAGCAAC
 8401 ACTCGGCTTG CCCCTATTCA AGCTGTAGCC CTGGGTCATC CTGCCACTAC
 8451 GCATTCTGAA TTTATTTGATT ATGTCATCGT AGAAGATGAT TATGTGGGCA
 8501 GTGAAGATTG TTTCAGCGAA ACCCTTTTAC GCTTACCCAA AGATGCCCTA
 8551 CCTTATGTAC CTTCTGCACT CGCCCCACAA AAAGTGGATT ATGTACTCAG
 8601 GGAAAACCCCT GAAGTAGTCA ATATCGGTAT TGCCGCTACC ACAATGAAAT
 8651 TAAACCCCTGA ATTTTGTCTA ACATTGCAAG AAATCAGAGA TAAAGCTAAA
 8701 GTCAAAATAC ATTTTCATTT CGCACTTGA CAATCAACAG GCTTGACACA
 8751 CCCTTATGTC AAATGGTTTA TCGAAAGCTA TTTAGGTGAC GATGCCACTG
 8801 CACATCCCCA CGCACCTTAT CACGATTATC TGGCAATATT GCGTGATTGC
 8851 GATATGCTAC TAAATCCGTT TCCTTTTCGGT AATACTAACG GCATAATTGA
 8901 TATGGTTACA TTAGGTTTAG TTGGTGTATG CAAAACGGGG GATGAAGTAC
 8951 ATGAACATAT TGATGAAGGT CTGTTTAAAC GCTTAGGACT ACCAGAAATGG
 9001 CTGATAGCCG ACACACGAGA AACATATATT GAATGTGCTT TGCGTCTAGC
 9051 AGAAAACCAT CAAGAACGCC TTGAACTCCG TCGTTACATC ATAGAAAACA

FIG. 7L.

9101 ACGGCTTACA AAAGCTTTT ACAGGCGACC CTCGTCCATT GGGCAAAATA
9151 CTGCTTAAGA AACAAATGA ATGGAAGCGG AAGCACTTGA GTAAAAAATA
9201 ACGGTTTTTT AAAGTAAAG TCGGGTTAAT TTTCAAAGCG TTTTAAAAAC
9251 CTCCTCAAAA TCAACCGCAC TTTTATCTTT ATAACGATCC CGCACGCTGA
9301 CAGTTTATCA GCCTCCCGCC ATAAAACTCC GCCTTTCATG GCGGAGATTT
9351 TAGCCAAAAC TGGCAGAAAT TAAAGGCTAA AATCACCAAA TTGCACCACA
9401 AAATCACCAA TA₂CCACAAA AAA

44 / 68

FIG. 8A.

1	GATCAATCTG	GCGATATTT	TTGCCAAAGG	TGGTAACATT	AATGTCCGCG
51	CTGCCACTAT	TCGCAATAAA	GGTAAACTTT	CTGCCGACTC	TGTAAGCAAA
101	GATAAAAGTG	GTAACATTGT	TCTCTCTGCC	AAAGAAAGTG	AAGCGGAAAT
151	TGGCGGTGTA	ATTTCCGCTC	AAATCAGCA	AGCCAAAGGT	GGTAAGTTGA
201	TGATTACAGG	CGATAAAGTT	ACATTGAAAA	CGGGTGCAGT	TATCGACCTT
251	TCGGGTAAAG	AAGGGGAGA	AACTTATCTT	GGCGGTGACG	AGCGTGGCGA
301	AGGTAAAAAC	GGCATTCAAT	TAGCAAAAGAA	AACCACTTTA	GAAAAAGGCT
351	CAACAATTAA	TGTGTCAGGT	AAAGAAAAAG	GTGGGCGCGC	TATTGTATGG
401	GGCGATATTG	CGTTAATTGA	CGGCAATATT	AATGCCCAAG	GTAAGATAT
451	CGCTAAAAC	GGTGGTTTGG	TGGAGACGTC	GGGGCATTAC	TTATCCATTG
501	ATGATAACGC	AATTGTTAAA	ACAAAAGAAT	GGCTACTAGA	CCCAGAGAAT
551	GTGACTATTG	AAGCTCCCTC	CGCTTCTCGC	GTCGAGCTGG	GTGCCGATAG
601	GAATTCCCCAC	TCGGCAGAGG	TGATAAAAGT	GACCCCTAAA	AAAAATAACA
651	CCTCCTTGAC	AACACTAACC	AATACAACCA	TTTCAAAATCT	TCTGAAAAGT
701	GCCCACGTGG	TGAACATAAC	GGCAAGGAGA	AAACTTACCG	TTAATAGCTC
751	TATCAGTATA	GAAAGAGGCT	CCCCTTAAT	TCTCCACAGT	GAAGGTCAGG

FIG. 8B.

801 GCGGTCAAGG TGTTCAGATT GATAAAGATA TTACTTCTGA AGGCGGAAAT
851 TTAACCATTT ATTCTGGCGG ATGGGTTGAT GTTCATAAAA ATATTACGCT
901 TGGTAGCGGC TTTTAAACA TCACAACCTAA AGAAGGAGAT ATCGCCTTCG
951 AAGACAAGTC TGGACGGAAC AACCTAACCA TTACAGCCCA AGGACCATC
1001 ACCTCAGGTA ATAGTAACGG CTTTAGATTT AACACGTCT CTCTAAACAG
1051 CCTTGGCGGA AAGCTGAGCT TTACTGACAG CAGAGAGGAC AGAGGTAGAA
1101 GAACTAAGG TAAATATCTCA AACAAATTG ACGGAACGTT AAACATTTCC
1151 GGAACCTGTAG ATATCTCAAT GAAAGCACCC AAAGTCAGCT GGTTTACAG
1201 AGACAAAGGA CGCACCTACT GGAACGTAAC CACTTTAAAT GTTACCTCGG
1251 GTAGTAAATT TAACCTCTCC ATTGACAGCA CAGGAAGTGG CTCAACAGGT
1301 CCAAGCATAC GCAATGCAGA ATTAATGGC ATAACATTTA ATAAAGCCAC
1351 TTTTAAATATC GCACAAGGCT CAACAGCTAA CTTTAGCATC AAGGCATCAA
1401 TAATGCCCTT TAAGAGTAAC GCTAACTACG CATTATTTAA TGAAGATATT
1451 TCAGTCTCAG GGGGGGGTAG CGTTAATTTC AAACCTTAACG CCTCATCTAG
1501 CAACATACAA ACCCCTGGCG TAAATTATAAA ATCTCAAAAC TTTAATGTCT
1551 CAGGAGGGTC AACTTTAAAT CTCGAAGGCTG AAGGTTCAAC AGAAACCGCT
1601 TTTTCAATAG AAAATGATTT AAACCTTAAC GCCACCGGTG GCAATATAAC

46/ 68

47/68

FIG. 8C.

1651 AATCAGACAA GTCGAGGGTA CCGATTCACG CGTCAACAAA GGTGTCGCAG
1701 CCAAAAAAAA CATAACTTTT AAAGGGGGTA ATATCACCTT CGGCTCTCAA
1751 AAAGCCACAA CAGAAATCAA AGGCAATGTT ACCATCAATA AAAACACTAA
1801 CGCTACTCTT CGTGGTGCGA ATTTTGCCGA AAACAAATCG CCTTTAAATA
1851 TAGCAGGAAA TGTTATTAAAT AATGGCAACC TTACCACCTG CGGCTCCATT
1901 ATCAATATAG CCGGAAATCT TACTGTTTCA AAAGGCGCTA ACCTTCAAGC
1951 TATAACAAAT TACACTTTTA ATGTAGCCGG CTCATTTGAC AACAAATGGCG
2001 CTTCAAACAT TTCCATTGCC AGAGGAGGGG CTAATTTAA AGATATCAAT
2051 AACACCAGTA GCTTAAATAT TACCACCAAC TCTGATACCA CTTACCGCAC
2101 CATATATAAA GGCAATATAT CCAACAAATC AGGTGATTTG AATATTATTG
2151 ATAAAAAAG CGACGCTGAA ATCCAAATTG GCGGCAATAT CTCACAAAAA
2201 GAAGGCAATC TCACAATTTC TTCTGATAAA GTAAATATTA CCAATCAGAT
2251 AACAAATCAA GCAGGCGTTG AAGGGGGCG TTCTGATTCA AGTGAGGCAG
2301 AAAATGCTAA CCTAACTATT CAAACCAAAG AGTTAAAATT GGCAGGAGAC
2351 CTAAATATTT CAGGCTTTAA TAAAGCAGAA ATTACAGCTA AAATGGCAG
2401 TGATTTAACT ATTGGCAATG CTAGCGGTGG TAATGCTGAT GCTAAAAAAG

FIG. 8D.

2451	TGACTTTTGA	CAAGGTAAA	GATCAAAA	TCTCGACTGA	CGGTCACAAT
2501	GTAACACTAA	ATAGCGAAGT	GAAACGTCT	AATGGTAGTA	GCAATGCTGG
2551	TAATGATAAC	AGCACCGGTT	TAACCATTTC	CGCAAAAGAT	GTAACGGTAA
2601	ACAATAACGT	TACCTCCCAC	AAGACAATAA	ATATCTCTGC	CGCAGCAGGA
2651	AATGTAACAA	CCAAAGAAAG	CACAACTATC	AATGCAACCA	CAGGCAGCGT
2701	GGAAGTAACT	GCTCAAAATG	GTACAATTAA	AGGCAACATT	ACCTCGCAAA
2751	ATGTAACAGT	GACAGCAACA	GAAAATCTTG	TTACCACACA	GAATGCTGTC
2801	ATTAATGCAA	CCAGCGGCAC	AGTAAACATT	AGTACAAAAA	CAGGGGATAT
2851	TAAAGGTGGA	ATTGAATCAA	CTTCCGGTAA	TGTAAATATT	ACAGCGAGCG
2901	GCAATACACT	TAAGGTAAGT	AATATCACTG	GTCAGATGT	AACAGTAACA
2951	GCGGATGCAG	GAGCCTTGAC	AACTACAGCA	GGCTCAACCA	TTAGTGCGAC
3001	AACAGGCAAT	GCAAATATTA	CAACCAAAAC	AGGTGATATC	AACGGTAAAG
3051	TTGAATCCAG	CTCCGGCTCT	GTAACACTTG	TTGCAACTGG	AGCAACTCTT
3101	GCTGTAGGTA	ATATTTCAGG	TAACACTGTT	ACTATTACTG	CGGATAGCGG
3151	TAAATTAAAC	TCCACAGTAG	GTTCTACAAT	TAATGGGACT	AATAGTGTA
3201	CCACCTCAAG	CCAATCAGGC	GATATTGAAG	GTACAAATTC	TGGTAATACA
3251	GTAAATGTTA	CAGCAAGCAC	TGGTGATTTA	ACTATTGGAA	ATAGTGCAAA

FIG. 8E.

3301 AGTTGAAGCG AAAAATGGAG CTGCAACCTT AACTGCTGAA TCAGGCAAAT
3351 TAACCACCCA AACAGGCTCT AGCATTACCT CAAGCAATGG TCAGACAACT
3401 CTTACAGCCA AGGATAGCAG TATCGCAGGA AACATTAAATG CTGCTAATGT
3451 GACGTTAAAT ACCACAGGCA CTTTAACTAC TACAGGGGAT TCAAAGATTA
3501 ACGCAACCAG TGGTACCTTA ACAATCAATG CAAAAGATGC CAAATTAGAT
3551 GGTGCTGCAT CAGGTGACCG CACAGTAGTA AATGCAACTA ACGCAAGTGG
3601 CTCCTGGTAAC GTGACTGCCA AAACCTCAAG CAGCGTGAAAT ATCACC GGGG
3651 ATTTAAACAC AATAAATGGG TTAAATATCA TTTTCGGAAAA TGGTAGAAAC
3701 ACTGTGCGCT TAAGAGGCAA GGAAATTGAT GTGAAATATA TCCAACCAGG
3751 TGTAGCAAGC GTAGAAGAGG TAATTGAAGC GAAACGCGTC CTGAGAAAGG
3801 TAAAAGATTT ATCTGATGAA GAAAGAGAAA CACTAGCCAA ACTTGGTGTA
3851 AGTGCTGTAC GTTTCGTTGA GCCAAATAAT GCCATTACGG TTAATACACA
3901 AAACGAGTTT ACAACCAAAC CATCAAGTCA AGTGACAATT TCTGAAGGTA
3951 AGGCGTGT TTCTCAAGTGGT AATGGCGCAC GAGTATGTAC CAATGTTGCT
4001 GACGATGGAC AGCAGTAGTC AGTAATTGAC AAGGTAGATT TCATCCTGCA
4051 ATGAAGTCAT TTTATTTTCG TATTATTAC TGTGTGGGTT AAAGTTCAGT

49/68

50/68

FIG. 8F.

4101 ACGGGCTTTA CCCACCTTGT AAAAAATTAC GAAAAATACA ATAAAGTATT
4151 TTAAACAGGT TATTATTATG AAAAACATAA AAAGCAGATT AAAACTCAGT
4201 GCAATATCAA TATTGCTTGG CTTGGCTTCT TCATCGACGT ATGCAGAAAG
4251 AGCGTTTTTA GTAAAGGCT TTCAGTTATC TGGCGCG

FIG. 9A.

1 GGAATGAGC GTCGTACACG GTACAGCAAC CATGCAAGTA GACGGCAATA
 51 AAACCACTAT CCGTAATAGC GTCAATGCTA TCATCAATTG GAAACAATTT
 101 AACATTGACC AAAATGAAAT GGAGCAGTTT TTACAAGAAA GCAGCAACTC
 151 TGCCGTTTTC AACCGTGTTA CATCTGACCA AATCTCCCAA TTAAAAGGGA
 201 TTTTAGATTC TAACGGACAA GTCTTTTAA TCAACCCAAA TGGTATCACA
 251 ATAGGTAAAG ACGCAATTAT TAACACTAAT GGCTTTACTG CTTCTACGCT
 301 AGACATTCTT AACGAAAACA TCAAGGCGCG TAATTTCCACC CTTGAGCAAA
 351 CCAAGGATAA AGCACTCGCT GAAATCGTGA ATCACGGTTT AATTACCGTT
 401 GGTAAGACG GTAGCGTAAA CCTTATTGGT GGCAAGTGA AAAACGAGGG
 451 CGTGATTAGC GTAAATGGCG GTAGTATTTC TTTACTTGCA GGCACAAAAA
 501 TCACCATCAG CGATATAATA AATCCAACCA TCACTTACAG CATTGCTGCA
 551 CCTGAAAACG AAGCGATCAA TCTGGGCGAT ATTTTGTCCA AAGTGTGTAA
 601 CATTAATGTC CGCGCTGCCA CTATTCGCAA TAAAGGTAAA CTTTCTGCCG
 651 ACTCTGTAG CAAAGATAAA AGTGGTAACA TTGTTCTCTC TGCCAAAGAA
 701 GGTGAAGCGG AAATTGGCGG TGTAAATTTC GCTCAAAATC AGCAAGCCAA
 751 AGGTGGTAAG TTGATGATTA CAGGTGATAA AGTCACATTA AAAACAGGTG

51 / 68

FIG. 9B.

801 CAGTTATCGA CCTTTCAGGT AAAGAAAGGG GAGAGACTTA TCCTGGCGGT
 851 GATGAGCGTG GCGAAGGTAA AAATGGTATT CAATTAGCGA AGAAAAACCTC
 901 TTTAGAAAAA GGCTCGACAA TTAATGTATC AGGCAAAGAA AAAGGCGGGC
 951 GCGCTATTGT ATGGGGCGGAT ATTGCATTAA TTAATGGTAA CATTAATGCT
 1001 CAAGGTAGCG ATATTGCTAA AACTGGCGGC TTTGTGGAAA CATCAGGACA
 1051 TGACTTATCC ATTGGTGATG ATGTGATTGT TGACGCTAAA GAGTGGTTAT
 1101 TAGACCCAGA TGATGTGTCC ATTGAAACTC TTACATCTGG ACGCAATAAT
 1151 ACCGCGGAAA ACCAAGGATA TACAACAGGA GATGGGACTA AAGAGTCACC
 1201 TAAAGGTAAT AGTATTTCTA AACCTACATT AACAAACTCA ACTCTTGAGC
 1251 AAATCCTAAG AAGAGGTTCT TATGTTAATA TCACTGCTAA TAATAGAAAT
 1301 TATGTTAATA GCTCCATCAA CTTATCTAAT GGCAGTTTAA CACTTCACAC
 1351 TAAACGAGAT GGAGTTAAAA TTAACGGTGA TATTACCTCA AACGAAAATG
 1401 GTAAATTTAAC CATTAAAGCA GGCTCTTGGG TTGATGTTCA TAAAAACATC
 1451 ACGCTTGGTA CGGGTTTTTT GAATATGTTC GCTGGGGATT CTGTAGCTTT
 1501 TGAGAGAGAG GCGGATAAAG CACGTAACGC AACAGATGCT CAAATTACCG
 1551 CACAAGGGAC GATAACCGTC AATAAAGATG ATAAACAATT TAGATTCAAT
 1601 AATGTATCTA TTAACGGGAC GGGCAAGGGT TTAAAGTTTA TTGCAAATCA

52 / 60

FIG. 9C.

1651 AAATAATTTC ACTCATAAAT TTGATGGCGA AATTAACATA TCTGGAATAG
 1701 TAACAATTAA CCAAAACCACG AAAAAAGATG TTAAATACTG GAATGCATCA
 1751 AAAGACTCTT ACTGGAATGT TTCTTCTCTT ACTTTGAATA CGGTGCAAAA
 1801 ATTTACCCTTT ATAAAATTCTG TTGATAGCGG CTCAAAATTCC CAAGATTGA
 1851 GGTCATCACG TAGAAGTTTT GCAGGCGTAC ATTTTAACGG CATCGGAGGC
 1901 AAAACAAACT TCAACATCGG AGCTAACGCA AAAGCCTTAT TTAAATTAAA
 1951 ACCAAACGCC GCTACAGACC CAAAAAAGA ATTACCTATT ACTTTTAACG
 2001 CCAACATTAC AGCTACCGGT AACAGTGATA GCTCTGTGAT GTTTGACATA
 2051 CACGCCAATC TTACCTCTAG AGCTGCCGGC ATAAACATGG ATTCAATTAA
 2101 CATTACCGGC GGGCTTGACT TTTCCATAAC ATCCCATAAAT CGCAATAGTA
 2151 ATGCTTTTGA AATCAAAAAA GACTTAACTA TAAATGCAAC TGGCTCGAAT
 2201 TTTAGTCTTA AGCAAACGAA AGATTCTTTT TATAATGAAT ACAGCAAACA
 2251 CGCCATTAACTCAAGTCATA ATCTAACCAT TCTTGGCGC AATGTCACTC
 2301 TAGGTGGGGA AAATTCAAGC AGTAGCATTA CGGGCAATAT CAATATCACC
 2351 AATAAAGCAA ATGTTACATT ACAAGCTGAC ACCAGCAACA GCAACACAGG
 2401 CTTGAAGAAA AGAACTCTAA CTCTTGGCAA TATATCTGTT GAGGGGAATT

53/88

FIG. 9D.

2451 TAAGCCCTAAC TGGTGCAAAT GCAAACATTG TCGGCAATCT TTCTATTGCA
 2501 GAAGATTCCA CATTTAAAGG AGAAGCCAGT GACAACCTAA ACATCACCGG
 2551 CACCTTTACC AACAAACGGTA CCGCCAACAT TAATATAAAA CAAGGAGTGG
 2601 TAAAACTCCA AGGCGATATT ATCAATAAAG GTGGTTTAAA TATCACTACT
 2651 AACGCCTCAG GCACTCAAAA AACCATTTAT AACGGAAATA TAACTAACGA
 2701 AAAAGGCGAC TTAAACATCA AGAATATTAA AGCCGACGCC GAAATCCAAA
 2751 TTGGCGGCAA TATCTCACAA AAAGAAGGCA ATCTCACAAAT TTCTTTCTGAT
 2801 AAAGTAAATA TTACCAATCA GATAACAATC AAAGCAGGCG TTGAAGGGGG
 2851 GCGTTCTGAT TCAAGTGAGG CAGAAAATGC TAACCTAACT ATTCAAACCA
 2901 AAGAGTTAAA ATTGGCAGGA GACCTAAATA TTTCAGGCTT TAATAAAGCA
 2951 GAAATTACAG CTAAAAAATGG CAGTGATTTA ACTATTGGCA ATGCTAGCCG
 3001 TGGTAATGCT GATGCTAAAA AAGTGACTTT TGACAAGGTT AAAGATTCAA
 3051 AAATCTCGAC TGACGGTCAC AATGTAACAC TAAATAGCGA AGTGAAAACG
 3101 TCTAATGGTA GTAGCAATGC TGGTAATGAT AACAGCACCG GTTTAACCAT
 3151 TTCCGCAAAA GATGTAACGG TAAACAATAA CGTTACCTCC CACAAGACAA
 3201 TAAATATCTC TGCCGCAGCA GGAAATGTAA CAACCAAGA AGGCACAACT
 3251 ATCAATGCAA CCACAGGCAG CGTGGAAGTA ACTGCTCAA ATGGTACAAT

54/68

FIG. 9E.

3301 TAAAGGCAAC ATTACCTCGC AAAATGTAAC AGTGACAGCA ACAGAAAATC
3351 TTGTTACCAC AGAGAATGCT GTCATTAATG CAACCAGCGG CACAGTAAAC
3401 ATTAGTACAA AAACAGGGA TATTAAAGGT GGAATTGAAT CAACTTCCGG
3451 TAATGTAAAT ATTACAGCGA GCGGCAATAC ACTTAAGGTA AGTAATATCA
3501 CTGGTCAAGA TGTAACAGTA ACAGCGGATG CAGGAGCCTT GACAACTACA
3551 GCAGGCTCAA CCATTAGTGC GACAACAGGC AATGCAAATA TTACAACCAA
3601 AACAGGTGAT ATCAACGGTA AAGTTGAATC CAGCTCCGGC TCTGTAACAC
3651 TTGTTGCAAC TGGAGCAACT CTTGCTGTAG GTAAATATTTC AGGTAACACT
3701 GTTACTATTA CTGCGGATAG CGGTAAATTA ACCTCCACAG TAGGTTCTAC
3751 AATTAATGGG ACTAATAGTG TAACCACCTC AAGCCAATCA GCGATATTG
3801 AAGGTACAAT TTCTGGTAAT ACAGTAAATG TTACAGCAAG CACTGGTGAT
3851 TTAACATATTG GAAATAGTGC AAAAGTTGAA GCGAAAATG GAGCTGCAAC
3901 CTTAACTGCT GAATCAGGCA AATTAACCAC CCAAACAGGC TCTAGCATTA
3951 CCTCAAGCAA TGGTCAGACA ACTCTTACAG CCAAGGATAG CAGTATCGCA
4001 GGAAACATTA ATGCTGCTAA TGTGACGTTA AATACCACAG GCACTTTAAC
4051 TACTACAGGG GATTCAAAGA TTAACGCAAC CAGTGGTACC TTAACAATCA

FIG. 9F.

4101 ATGCAAAAGA TGCCAAATTA GATGGTGCTG CATCAGGTGA CCGCACAGTA
4151 GTAAATGCAA CTAACGCAAG TGGCTCTGGT AACGTGACTG CGAAAACCTC
4201 AAGCAGCGTG AATATCACCG GGGATTAAA CACAATAAAT GGGTAAATA
4251 TCATTTCCGA AAATGGTAGA AACACTGTGC GCTTAAGAGG CAAGGAAATT
4301 GATGTGAAT ATATCCAACC AGGTGTAGCA AGCGTAGAAG AGGTAATTGA
4351 AGCGAAACGC GTCCTTGAGA AGGTAAAAGA TTTATCTGAT GAAGAAAGAG
4401 AAACACTAGC CAAACTTGGT GTAAGTGCTG TACGTTTCGT TGAGCCAAAT
4451 AATGCCATTA CGGTTAATAC ACAAACGAG TTTACAACCA AACCATCAAG
4501 TCAAGTGACA ATTTCTGAAG GTAAGGCCGTG TTTCTCAAGT GGTAATGGCG
4551 CACGAGTAG TACCAATGTT GCTGACGATG GACAGCAGTA GTCAGTAATT
4601 GACAAGGTAG ATTTCATCCT GCAATGAAGT CATTTTATT TCGTATTATT
4651 TACTGTGTGG GTTAAAGTTC AGTACGGGCT TTACCCACCT TGTAATAAAT
4701 TA

56/68

FIG. 10A. COMPARISON OF DERIVED AMINO ACID SEQUENCE

	1	50	
Hmw3com
Hmw4com
Hmw1com	MNKIYRLKFS	KRLNALVAVS	ELARGCDHST EKGSEKPARM KVRHLALKPL
Hmw2com	MNKIYRLKFS	KRLNALVAVS	ELARGCDHST EKGSEKPARM KVRHLALKPL
	51	57/68	100
Hmw3com
Hmw4com
Hmw1com	SAMLLSLGVT	SIPQSVLASG	LQMSVVHGT ATMQVDGNKT TIRNSVNAII
Hmw2com	SAMLLSLGVT	SIPQSVLASG	LQMSVVHGT ATMQVDGNKT TIRNSVNAII
	101		150
Hmw3com
Hmw4com	NWKQFNIDQN	EMEQFLQESS	NSAVFNRTS DQISQLKGIL DSNQGVFLIN

FIG. 10B.

Hmw1com	NWKQFNIDQN	EMVQFLQENN	NSAVFN RVTS	NQISQLKGIL	DSNGQVFLIN	
Hmw2com	NWKQFNIDQN	EMVQFLQENN	NSAVFN RVTS	NQISQLKGIL	DSNGQVFLIN	
						200
Hmw3com	
Hmw4com	PNGITIGKDA	IINTNGFTAS	TLDISNENIK	ARNFTLEQTK	DKALAEIVNH	
Hmw1com	PNGITIGKDA	IINTNGFTAS	TLDISNENIK	ARNFTLEQTK	DKALAEIVNH	58/88
Hmw2com	PNGITIGKDA	IINTNGFTAS	TLDISNENIK	ARNFTLEQTK	DKALAEIVNH	88
						250
Hmw3com	
Hmw4com	GLITVGKDGS	VNLIGGKVKN	EGVISVNGGS	ISLLAGQKIT	ISDIINPTIT	
Hmw1com	GLITVGKDGS	VNLIGGKVKN	EGVISVNGGS	ISLLAGQKIT	ISDIINPTIT	
Hmw2com	GLITVGKDGS	VNLIGGKVKN	EGVISVNGGS	ISLLAGQKIT	ISDIINPTIT	
						300
Hmw3com	INLGDIFAKG	GNINVRAATI	RNKGKLSADS	VSKDKSGNIV	
						251

FIG. 10C.

Hmw4com	YSIAAPENEA	INLGDIFAKG	GNINVRAATI	RNKGKLSADS	VSKDKSGNIV
Hmw1com	YSIAAPENEA	VNLGDIFAKG	GNINVRAATI	RNKGKLSADS	VSKDKSGNIV
Hmw2com	YSIAAPENEA	VNLGDIFAKG	GNINVRAATI	RNKGKLSADS	VSKDKSGNIV

350

Hmw3com	LSAKEGEAEI	GGVISAQNQQ	AKGGKLMITG	DKVTLKTGAV	IDLSGKEGGE
Hmw4com	LSAKEGEAEI	GGVISAQNQQ	AKGGKLMITG	DKVTLKTGAV	IDLSGKEGGE
Hmw1com	LSAKEGEAEI	GGVISAQNQQ	AKGGKLMITG	DKVTLKTGAV	IDLSGKEGGE
Hmw2com	LSAKEGEAEI	GGVISAQNQQ	AKGGKLMITG	DKVTLKTGAV	IDLSGKEGGE

59/68

400

351

Hmw3com	TYLGGDERGE	GKNGIQLAKK	TITLEKGSTIN	VSGKEKGGA	IVWGDIALID
Hmw4com	TYLGGDERGE	GKNGIQLAKK	TITLEKGSTIN	VSGKEKGGA	IVWGDIALID
Hmw1com	TYLGGDERGE	GKNGIQLAKK	TITLEKGSTIN	VSGKEKGGA	IVWGDIALID
Hmw2com	TYLGGDERGE	GKNGIQLAKK	TITLEKGSTIN	VSGKEKGGA	IVWGDIALID

FIG. 10D.

						450
Hmw3com	GNINAQK.D	IAKTGGFVET	SGHYLSIDDN	AIVKTKEWLL	DPENVTIEAP	
Hmw4com	GNINAQGS.D	IAKTGGFVET	SGHDLSIGDD	VIVDAKEWLL	DPDDVSIETL	
Hmw1com	GNINAQGS.D	IAKTGGFVET	SGHDLFIKDN	AIVDAKEWLL	DPDNVTINAE	
Hmw2com	GNINAQGS.D	IAKTGGFVET	SGHYLSIESN	AIVKTKEWLL	DPDDVTIEAE	
						500
Hmw3com	SASRVELGAD	RNSHSAEVIK	VTLKKNNTSL	TTLTNTTISN	LLKSAHVNI	
Hmw4com	TSGRNNTGEN	QGYTTGDGTK	ESPKGNSISK	PTLTNSTLEQ	ILRRGSYVNI	
Hmw1com	TACRSNTSED	DEYTGSGNSA	STPKRNKE.K	TTLTNTTLES	ILKKGTFVNI	
Hmw2com	DPLRNNTGIN	DEFPTGTGEA	SDPKKNSELK	TTLTNTTISN	YLKNAWTMNI	
						550
Hmw3com	TARRKLTVNS	SISIERGSHL	ILHSEGQGGQ	GVQIDKDITS	.E...GGNLT	
Hmw4com	TANNRIYVNS	SINLSNGS.L	TLHTK...RD	GVKINGDITS	NE...NGNLT	
Hmw1com	TANQRIYVNS	SINL.SNGSL	TLWSEGRSGG	GVEINNNDITT	GDDTRGANLT	
Hmw2com	TASRKLTVNS	SINGSNGLSHL	ILHSGQRGG	GVQIDGDIT.	...SKGGNLT	

FIG. 10E.

	551		600
Hmw3com	IYSGGWVDVH	KNITLGS.GF	LNITKEGDI AFEDKSGR... ..NNLTITAQ
Hmw4com	IKAGSWVDVH	KNITLGT.GF	LNIVAGDS.V AFEREGDKAR NATDAQITAQ
Hmw1com	IYSGGWVDVH	KNISLGAQGN	INITAKQD.I AFEKGSNQV.ITGQ
Hmw2com	IYSGGWVDVH	KNITLD.QGF	LNITA.AS.V AFEGGNNKAR DANNLTITAQ
	601		650
Hmw3com	GTITSG.NSN	GFRFNNVSLN	SLGGKLSFTD SREDRGRRTK GNISNKFDDGT
Hmw4com	GTITVKNKDDK	QFRFNNVSN	GTGKGLKFIA NQN..... .NFTHKFDGE
Hmw1com	GTIT.SGNQK	GFRFNNVSLN	GTGSGLQFTT KRTN.....K YAITNKFEGT
Hmw2com	GTVTITGEGK	DFRANNVSLN	GTGKGLNIIS SVNN..... .LTHNLSGT
	651		700
Hmw3com	LNISGTVDIS	MKAPKVSIFY	RD.KGRTYWN VTTLNVTSGS KFNLSIDSTG
Hmw4com	INISGIVTIN	QTTKKDKVKYW	NA.SKDSYWN VSSLTLNTVQ KFTF.IKFVD
Hmw1com	LNISGKVNIS	MVLPKNESGY	DKFKGRTYWN LTSLNVSESG EFNLTIDSRG

FIG. 10F.

Hmw2com INISGNITIN QTRKNTSYW QTSHD.SHWN VSALNLETGA NFTF.IKYIS

701 750

Hmw3com SGSTG...PS IRNA..ELNG ITFN...KA TFNIAQGSTA NFSIKASIMP

Hmw4com SGSNS...QD LRSSRRSFAG VHFNGIGGKT NFNIGANAKA LFKLKPNAAT

Hmw1com SDSAGTLTQ.PYNLNG ISFN...KDT TFNVERNARV NFDIKAPIGI

Hmw2com SNSKGLTTQY RSSAGVNFNG V..N...GNM SFNLKEGAKV NFKLKPENNM 62/68

751 800

Hmw3com FKSANANYAL. FNEDISVSG. .GGSVNFKLN ASSSNIQTPG VIKSQNFNV

Hmw4com DPKKELPIT. FNANITATGN SDSSVMFDIH A...NLTSRA AGINMDSINI

Hmw1com NKYSSLNyas FNGNISVSG. .GGSVDFTL ASSSNVQTPG VVINSKYFNV

Hmw2com NTSKPLPI.R FLANITATG. .GGSVFFDIY ANHS...GRG AFLKMSEINI

801 850

Hmw3com SGGSTLNLKA EGSTETAFSI ENDLNLNATG GNITIRQVEG T..DSRVNKG

Hmw4com TGGLDFSITS HNRNSNAFEI KKDLTINATG SNFSLKQTKD SFYNEYSKHA

FIG. 10G.

Hmw1com	STGSSLRFKT	SGSTKTGFSI	EKDLTLNATG	GNITLLQVEG	T..DGMIGKG	
Hmw2com	SNGANFTLNS	HVRGDDAFKI	NKDLTINATN	SNFSLRQTKD	DFYDGYARNA	
						851 900
Hmw3com	VAAKKNITFK	GGNITFGSQK	ATTEIKGNVT	INKNTNATLR	GANFAEN...	
Hmw4com	INSSHNLTIL	GGNVTLGGEN	SSSITGNIN	ITNKANVTLQ	ADTSNSNTGL	63/68
Hmw1com	IVAKKNITFE	GGNITFGSRK	AVTEIEGNVT	INNANAVTLI	GSDFDNHQ..	
Hmw2com	INSTYNISIL	GGNVTLGGQN	SSSITGNIT	IEKAANVTLE	ANNAPNQQNI	
						901 950
Hmw3com	KSPLNIAGNV	INNGNLTTAG	SIINIAGNLT	VSKGANLQAI	TNYTFNVAGS	
Hmw4com	KKRTLTLGNI	SVEGNLSLTG	ANANIVGNLS	IAEDSTFKGE	ASDNLNITGT	
Hmw1com	KPLTIKKDVI	INSGNLTAGG	NIVNIAGNLT	VESNANFKAI	TNFTFNVGGL	
Hmw2com	RDRVIKLGSL	LVNGSLSLTG	ENADIKGNLT	ISESATFKGK	TRDTLNTGN	
						951 1000

FIG. 10H.

Hmw3com	FDNNGASNIS	IARGGAKFK.	DINNTSSLNI	TTNSDTTYRT	IIKGNISNKS
Hmw4com	FTNNGTANIN	IKQGVVKLQG	DINNKGGLNI	TTNASGTQKT	IINGNITNEK
Hmw1com	FDNKGNSNIS	IAKGGARFK.	DIDNSKNLSI	TTNSSSTYRT	IISGNITNKN
Hmw2com	FTNNGTAEIN	ITQGVVKLG.	NVTNDGDLNI	TTHAKRNQRS	IIGGDIINNK

1001 1050
64/68

Hmw3com	GDLNIIDKKS	DAEIQIGGNI	SQKEGNLTIS	SDKVNITNQI	TIKAGVEGGR
Hmw4com	GDLNIKNIKA	DAEIQIGGNI	SQKEGNLTIS	SDKVNITNQI	TIKAGVEGGR
Hmw1com	GDLNITNEGS	DTEMQIGGDI	SQKEGNLTIS	SDKINITKQI	TIKAGVDGEN
Hmw2com	GSLNITDSNN	DAEIQIGGNI	SQKEGNLTIS	SDKINITKQI	TIKKGIDGED

1100

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Hmw3com	SDSSEAENAN	LTIQTKELKL	AGDLNISGFN	KAEITAKNGS	DLTIGNASGG
Hmw4com	SDSSEAENAN	LTIQTKELKL	AGDLNISGFN	KAEITAKNGS	DLTIGNASGG
Hmw1com	SDSDATNNAN	LTIKTKELKL	TQDLNISGFN	KAEITAKDGS	DLTIGNTNSA
Hmw2com	SSSDATSNAN	LTIKTKELKL	TEDLSISGFN	KAEITAKDGR	DLTIGNSNDG

FIG. 10I.

	1101	1150
Hmw3com	N..ADAKKVT FDKVKDSKIS TDGHNVTLNS EVKT..SNGS	SNAGNDNSTG
Hmw4com	N..ADAKKVT FDKVKDSKIS TDGHNVTLNS EVKT..SNGS	SNAGNDNSTG
Hmw1com	D.GTNAKKVT FNQVKDSKIS ADGHKVTLHS KVETSGSNNN	TEDSSDNNAG
Hmw2com	NSGAEAKKVT FNNVKDSKIS ADGHNVTLNS KVKTSSSNGG	RESNSDNDTG
	1151	1200
Hmw3com	LTISAKDVTV NNNVTSHKTI NISAAAGNVT TKEGTTINAT	TGSVEVTAQN
Hmw4com	LTISAKDVTV NNNVTSHKTI NISAAAGNVT TKEGTTINAT	TGSVEVTAQN
Hmw1com	LTIDAKNVTV NNNITSHKAV SISATSGEIT TKTGTTINAT	TGNVEIT...
Hmw2com	LTITAKNVEV NKDVTSLKTV NITA.SEKVT TTAGSTINAT	NGKASIT...
	1201	1250
Hmw3com	GTIKGNITSQ NVTVTATENL VTENAVINA TSGTVNISTK	TGDIKGGIES
Hmw4com	GTIKGNITSQ NVTVTATENL VTENAVINA TSGTVNISTK	TGDIKGGIES
Hmw1comAQ TGDIKGGIES

FIG. 10J.

Hmw2com	TK T.....	
					1300
Hmw3com	TSGNVNITAS	GNTLKVSNIT	GQDVTVTADA	GALT'TTAGST	ISATTGNANI
Hmw4com	TSGNVNITAS	GNTLKVSNIT	GQDVTVTADA	GALT'TTAGST	ISATTGNANI
Hmw1com	SSGSVTLTAT	EGALAVSNIS	GNTVTVTANS	GALT'TLAGST	IKG.TESVTT
Hmw2com	66 / 68
					1350
					1301
Hmw3com	TTKTGDINGK	VESSSGSVTL	VATGATLAVG	NISGNTVTIT	ADSGKLTSTV
Hmw4com	TTKTGDINGK	VESSSGSVTL	VATGATLAVG	NISGNTVTIT	ADSGKLTSTV
Hmw1com	SSQSGDIG..G	TISGGTVEVK	ATESLTTQSN
Hmw2comGDIS..G	TISGNTVSVS	ATVDLTTKSG
					1400
					1351
Hmw3com	GSTINGTNSV	TTSSQSGDIE	GTISGNTVNV	TASTGDLTIG	NSAKVEAKNG
Hmw4com	GSTINGTNSV	TTSSQSGDIE	GTISGNTVNV	TASTGDLTIG	NSAKVEAKNG

FIG. 10K.

Hmw1com SKIKATTGEA NVTSAATGTIG GTISGNTVNV TANAGDLTVG NGAEGINATEG
Hmw2com SKIEAKSGEA NVTSAATGTIG GTISGNTVNV TANAGDLTVG NGAEGINATEG

1401 1450

Hmw3com AATLTAESGK LTTQTGSSIT SSNGQTTLTA KDSSIAGNIN AANVTLNNTTG
Hmw4com AATLTAESGK LTTQTGSSIT SSNGQTTLTA KDSSIAGNIN AANVTLNNTTG
Hmw1com AATLTTSSGK LTTEASSHIT SAKGQVNLSA QDSSVAGSIN AANVTLNNTTG
Hmw2com AATLTATGNT LTTEAGSSIT STKGQVDLLA QNSSIAGNIN AANVTLNNTTG

67 / 68

1451 1500

Hmw3com TLTTTGDSKI NATSGTLTIN AKDAKLDGAA SGDRTVVNAT NASGSGNVTA
Hmw4com TLTTTGDSKI NATSGTLTIN AKDAKLDGAA SGDRTVVNAT NASGSGNVTA
Hmw1com TLTTVKGSNI NATSGTLTIN AKDAELNGAA LGNHTVVNAT NANGSGSVIA
Hmw2com TLTTVAGSDI KATSGTLTIN AKDAKLNGBA SGDSTEVENAV NASGSGSVTA

1501 1550

FIG. 10L.

Hmw3com	KTSSSVNITG	DLNTINGLNI	ISENGRNTVR	LRGKEIDVKY	IQPGVASVEE
Hmw4com	KTSSSVNITG	DLNTINGLNI	ISENGRNTVR	LRGKEIDVKY	IQPGVASVEE
Hmw1com	TTSSRVNITG	DLITINGLNI	ISKNGINTVL	LKGVKIDVKY	IQPGIASVDE
Hmw2com	ATSSSVNITG	DLNTVINGLNI	ISKDGRNTVR	LRGKEIEVKY	IQPGVASVEE
	1551			1600	
Hmw3com	VIEAKRVLEK	VKDLSDEERE	TLAKLGVSAV	RFVEPNNAIT	VNTQNEFTTK
Hmw4com	VIEAKRVLEK	VKDLSDEERE	TLAKLGVSAV	RFVEPNNAIT	VNTQNEFTTK
Hmw1com	VIEAKRILEK	VKDLSDEERE	ALAKLGVSAV	RFIEPNNTIT	VDTQNEFATR
Hmw2com	VIEAKRVLEK	VKDLSDEERE	TLAKLGVSAV	RFVEPNNTIT	VNTQNEFTTR
	1601			1632	
Hmw3com	PSSQVTISEG	KACFSSGNGA	RVCTNVADDG	QQ	
Hmw4com	PSSQVTISEG	KACFSSGNGA	RVCTNVADDG	QQ	
Hmw1com	PLSRIVISEG	RACFSNSDGA	TVCVNIADNG	R.	
Hmw2com	PSSQVIISEG	KACFSSGNGA	RVCTNVADDG	QP	

INTERNATIONAL SEARCH REPORT

International application No.

PCT/US93/02166

A. CLASSIFICATION OF SUBJECT MATTER

IPC(5) : C07K 13/00, 15/04, 17/02; C07H 21/04; C12N 15/09, 15/31; A61K 39/02

US CL : 530/350, 825; 536/27; 424/88, 92; 435/69.3

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

U.S. : 530/350, 825; 536/27; 424/88, 92; 435/69.3

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)

MEDLINE, APS, IG SUITE

search terms: high molecular weight protein, haemophilus

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Y,P	The Journal of Infectious Diseases, Volume 165(Suppl.), issued August 1992, S.J.Barenkamp., "Outer Membrane Protein and Lipopolysaccharides of Nontypeable <i>Haemophilus influenzae</i> ", pages S181-S184, see entire document.	1-19
Y,P	Infection and Immunity, Volume 60(4), issued April 1992, S.J.Barenkamp et al, "Cloning, Expression and DNA Sequence Analysis of Genes Encoding Nontypeable <i>Haemophilus influenzae</i> High-Molecular-Weight Surface-Exposed Proteins Related to Filamentous Hemagglutinin of <i>Bordetella pertussis</i> ", pages 1302-1313, see entire document.	1-19

☒ Further documents are listed in the continuation of Box C.
 ☐ See patent family annex.

* Special categories of cited documents:	*T	later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention	
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*E	earlier document published on or after the international filing date	*X	document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone
*L	document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)	*Y	document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art
*O	document referring to an oral disclosure, use, exhibition or other means		
*P	document published prior to the international filing date but later than the priority date claimed	*Z	document member of the same patent family

Date of the actual completion of the international search

14 May 1993

Date of mailing of the international search report

21 MAY 1993

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INTERNATIONAL SEARCH REPORT

International application No.
PCT/US93/02166

C (Continuation). DOCUMENTS CONSIDERED TO BE RELEVANT

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Y	Infection and Immunity, Volume 56(1), issued January 1988, E.J.Hansen, "Immune Enhancement of Pulmonary Clearance on Nontypable <i>Haemophilus influenzae</i> , pages 182-190, see entire document, especially Figures 3 and 4.	1-19
Y	Infection and Immunity, Volume 52(2), issued May 1986, S.J.Barenkamp, "Protection by Serum Antibodies in Experimental Nontypable <i>Haemophilus influenzae</i> Otitis Media", pages 572-578, see Figures 1 and 2.	1-19
Y	Proceedings of the National Academy of Sciences USA, Volume 80, issued March 1983, R.A.Young et al, "Efficient Isolation of Genes by Using Antibody Probes", pages 1194-1198, see entire document.	1-19
Y	Infection and Immunity, Volume 45(3), issued September 1984, R. Schneerson et al, "Serum Antibody Responses of Juvenile and Infant Rhesus Monkeys Injected with <i>Haemophilus influenzae</i> Type b and Pneumococcus Type 6A Capsular Polysaccharide-Protein Conjugates", pages 582-591, see entire document.	16-17
Y	Journal of Molecular Biology, Volume 157, issued 1982, J.Kyte et al, "A Simple Method for Displaying the Hydropathic Character of a Protein", pages 105-132, see entire document.	18-19
Y	Proceedings of the National Academy of Sciences, Volume 78(6), issued June 1981, T.P.Hopp et al, "Prediction of Protein Antigenic Determinants from Amino Acid Sequences", pages 3824-3828, see entire document.	18-19